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Substituted benzoylcyclohexanediones

Donna J. Veatch (Name of person realing paper or fee)

The invention relates to novel substituted benzoylcyclohexanediones, to processes for their preparation and to their use as herbicides.

It is already known that certain substituted benzoylcyclohexanediones have herbicidal properties (cf. EP-A-090262, EP-A-135191, EP-A-186118, EP-A-186119, EP-A-186120, EP-A-319075, WO-A-96/26200, WO-A-97/46530, WO-A-99/07688). However, the activity of these compounds is not in all respects satisfactory.

This invention, accordingly, provides the novel substituted benzoylcyclohexanediones of the general formula (I),

$$(R^{2})_{m} \xrightarrow{Q} Q \qquad (R^{4})_{n}$$

$$R^{1} \qquad Q \qquad (I)$$

in which 15

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- represents the numbers 0, 1, 2 or 3, m
- represents the numbers 0, 1, 2 or 3, n
- represents the single bond or represents alkanediyl (alkylene). A
- represents hydrogen or represents in each case optionally substituted alkyl or R^1 alkoxycarbonyl,
- represents optionally substituted alkyl, or together with R1 represents R^2 alkanediyl (alkylene) where in this case m represents 1 and R^1 and R^2 are

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located at the same carbon atom ("geminal") or at two adjacent carbon atoms ("vicinal"),

- R³ represents hydrogen, nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, halogen, or represents in each case optionally substituted alkyl, alkoxy, alkylthio, alkylsulphinyl, alkylsulphonyl, alkylamino, dialkylamino or dialkylaminosulphonyl,
- R4 represents nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, halogen, or represents in each case optionally substituted alkyl, alkoxy, alkylthio, alkylsulphinyl, alkylsulphonyl, alkylamino, dialkylamino or dialkylaminosulphonyl, and
- represents an optionally substituted 4- to 12-membered, saturated or unsaturated, monocyclic or bicyclic, heterocyclic grouping which contains 1 to 4 heteroatoms (up to 4 nitrogen atoms and, if appropriate, alternatively or additionally one oxygen atom or one sulphur atom, or one SO grouping or one SO₂ grouping), and which additionally contains one to three oxo groups (C=O) and/or thioxo groups (C=S) as components of the heterocycle,
 - including all possible tautomeric forms of the compounds of the general formula (I) and the possible salts of the compounds of the general formula (I).
- In the definitions, the hydrocarbon chains, such as alkyl or alkanediyl, are in each case straight-chain or branched including in combination with heteroatoms, such as in alkoxy.

In addition to the compounds of the general formula (I) - above - it is in each case also possible for the corresponding tautomeric forms - shown in exemplary manner below - to be present.

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$$(R^2)_m$$
 R^1
 $(R^4)_n$
 R^3

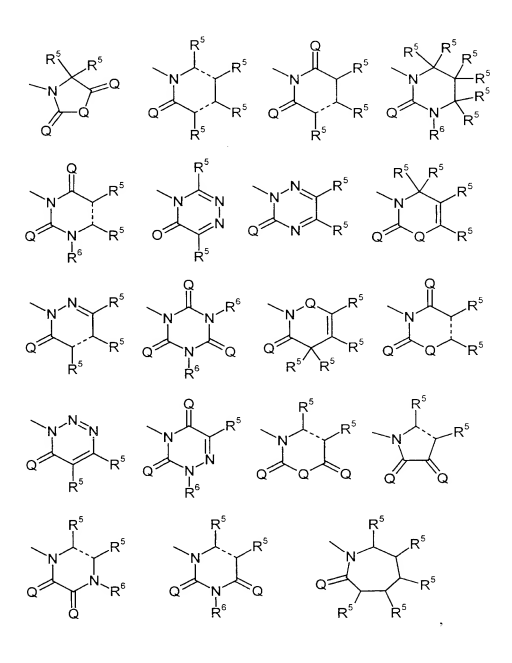
$$(R^2)_m$$
 $(R^4)_n$
 R^3

Preferred substituents of the radicals listed in the formula shown above are illustrated below:

- m preferably represents the numbers 0, 1 or 2.
- n preferably represents the numbers 0, 1 or 2.
- 10 A preferably represents a single bond or represents alkanediyl (alkylene) having 1 to 4 carbon atoms.
 - preferably represents hydrogen, represents optionally halogen-, C₁-C₄-alkoxy-, C₁-C₄-alkylthio-, C₁-C₄-alkylsulphinyl- or C₁-C₄-alkylsulphonyl-substituted alkyl having 1 to 6 carbon atoms or represents alkoxycarbonyl having up to 6 carbon atoms.
 - preferably represents optionally halogen-substituted alkyl having 1 to 6 carbon atoms, or together with R¹ represents alkanediyl (alkylene) having 2 to 5 carbon atoms, where in this case m represents 1 and R¹ and R² are located at the same carbon atom ("geminal") or at two adjacent carbon atoms ("vicinal").
- preferably represents hydrogen, nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, halogen, represents in each case optionally halogen-, C₁-C₄alkoxy-, C₁-C₄-alkylthio-, C₁-C₄-alkylsulphinyl- or C₁-C₄-alkylsulphonylsubstituted alkyl, alkoxy, alkylthio, alkylsulphinyl or alkylsulphonyl having

in each case up to 4 carbon atoms in the alkyl groups, or represents alkylamino, dialkylamino or dialkylaminosulphonyl having in each case up to 4 carbon atoms in the alkyl groups.

- preferably represents nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, halogen, represents in each case optionally halogen-, C₁-C₄-alkoxy-, C₁-C₄-alkylthio-, C₁-C₄-alkylsulphinyl- or C₁-C₄-alkylsulphonyl-substituted alkyl, alkoxy, alkylthio, alkylsulphinyl or alkylsulphonyl having in each case up to 4 carbon atoms in the alkyl groups, or represents alkylamino, dialkylamino or dialkylaminosulphonyl having in each case up to 4 carbon atoms in the alkyl groups.
 - Z preferably represents one of the heterocyclic groupings below



in which the bond drawn broken in each case denotes a single bond or a double bond,

Q represents oxygen or sulphur,

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R⁵ represents hydrogen, hydroxyl, mercapto, cyano, halogen, represents in each case optionally cyano-, halogen-, C₁-C₄-alkoxy-, C₁-C₄-

C₁-C₄-alkylsulphinyl- or C₁-C₄-alkylsulphonylalkylthio-, substituted alkyl, alkylcarbonyl, alkoxy, alkoxycarbonyl, alkylthio. alkylsulphinyl or alkylsulphonyl having in each case up to 6 carbon atoms in the alkyl groups, represents propadienylthio, represents in each case optionally halogen-substituted alkylamino or dialkylamino having in each case up to 6 carbon atoms in the alkyl groups, represents in each case optionally halogen-substituted alkenyl, alkinyl, alkenyloxy, alkenylthio or alkenylamino having in each case up to 6 carbon atoms in the alkenyl or alkinyl groups, represents in each case optionally halogen-substituted cycloalkyl, cycloalkyloxy, cycloalkylthio, cycloalkylamino, cycloalkylalkyl, cycloalkylalkoxy, cycloalkylalkylthio or cycloalkylalkylamino having in each case 3 to 6 carbon atoms in the cycloalkyl groups and optionally up to 4 carbon atoms in the alkyl moiety, or represents in each case optionally halogen-, C1-C4-alkyl- or C1-C4-alkoxy-substituted phenyl, phenyloxy, phenylthio, phenylamino, benzyl, benzyloxy, benzylthio or benzylamino, represents pyrrolidino, piperidino or morpholino, or - in the case that two adjacent radicals R5 and R5 are located at a double bond - together with the adjacent radical R^5 also represents a benzo grouping, and

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R⁶

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represents hydrogen, hydroxyl, amino, alkylideneamino having up to 4 carbon atoms, represents in each case optionally halogen- or C₁-C₄-alkoxy-substituted alkyl, alkoxy, alkylamino, dialkylamino or alkanoylamino having in each case up to 6 carbon atoms in the alkyl groups, represents in each case optionally halogen-substituted alkenyl, alkinyl or alkenyloxy having in each case up to 6 carbon atoms in the alkenyl or alkinyl groups, represents in each case optionally halogen-substituted cycloalkyl, cycloalkylalkyl or cycloalkylamino having in each case 3 to 6 carbon atoms in the cycloalkyl groups and optionally up to 3 carbon atoms in the alkyl moiety, or represents in each case

optionally halogen-, C_1 - C_4 -alkyl- or C_1 - C_4 -alkoxy-substituted phenyl or benzyl, or together with an adjacent radical R^5 or R^6 represents optionally halogen- or C_1 - C_4 -alkyl-substituted alkanediyl having 3 to 5 carbon atoms,

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where the individual radicals R⁵ and R⁶ - if two or more of them are attached to the same heterocyclic groupings, may have identical or different meanings in the context of the above definition.

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A particularly preferably represents a single bond, methylene, ethylidene (ethane-1,1-diyl) or dimethylene (ethane-1,2-diyl).

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R¹ particularly preferably represents hydrogen, represents in each case optionally fluorine-, chlorine-, methoxy-, ethoxy-, n- or i-propoxy-, methylthio-, ethylthio-, n- or i-propylthio-, methylsulphinyl-, ethylsulphinyl-, n- or i-propylsulphinyl-, methylsulphonyl-, ethylsulphonyl-, n- or i-propylsulphonyl-substituted methyl, ethyl, n- or i-propyl, n-, i- or s-butyl, or represents methoxycarbonyl, ethoxycarbonyl, n- or i-propoxycarbonyl.

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particularly preferably represents methyl, ethyl, n- or i-propyl, or together with R¹ represents methylene, ethane-1,1-diyl (ethylidene, -CH(CH₃)-), ethane-1,2-diyl (dimethylene, -CH₂CH₂-), propane-1,3-diyl (trimethylene, -CH₂CH₂CH₂-), butane-1,4-diyl (tetramethylene, -CH₂CH₂CH₂CH₂-) or pentane-1,5-diyl (pentamethylene, -CH₂CH₂CH₂CH₂CH₂-), where in this case m represents 1 and R¹ and R² are located at the same carbon atom ("geminal") or at two adjacent carbon atoms ("vicinal").

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particularly preferably represents hydrogen, nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, fluorine, chlorine, bromine, iodine, represents in each case optionally fluorine- and/or chlorine-, methoxy-, ethoxy-, n- or i-propoxy-, methylthio-, ethylthio-, n- or i-propylthio-, methylsulphinyl-.

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ethylsulphinyl-, methylsulphonyl- or ethylsulphonyl-substituted methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, represents in each case optionally fluorine- and/or chlorine-, methoxy-, ethoxy-, n- or i-propoxy-substituted methoxy, ethoxy, n- or i-propoxy, represents in each case optionally fluorine-and/or chlorine-substituted methylthio, ethylthio, n- or i-propylthio, methylsulphinyl, ethylsulphinyl, n- or i-propylsulphinyl, methylsulphonyl, ethylsulphonyl, n- or i-propylsulphonyl, or represents methylamino, ethylamino, n- or i-propylamino, dimethylamino, diethylamino, dimethylaminosulphonyl or diethylaminosulphonyl.

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 R^4

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particularly preferably represents nitro, cyano, carboxyl, carbamoyl, thiocarbamoyl, fluorine, chlorine, bromine, represents in each case optionally fluorine- and/or chlorine-, methoxy-, ethoxy-, n- or i-propoxy-, methylthio-, ethylthio-, n- or i-propylthio-, methylsulphinyl-, ethylsulphinyl-, methylsulphonyl- or ethylsulphonyl-substituted methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, represents in each case optionally fluorine- and/or chlorine-, methoxy-, ethoxy-, n- or i-propoxy-substituted methoxy, ethoxy, n- or i-propoxy, represents in each case optionally fluorine- and/or chlorine-substituted methylthio, ethylthio, n- or i-propylthio, methylsulphinyl, ethylsulphinyl, n- or i-propylsulphinyl, methylsulphonyl, ethylsulphonyl, n- or i-propylsulphonyl, or represents methylamino, ethylamino, n- or i-propylamino, dimethylamino, diethylamino, dimethylaminosulphonyl or diethylaminosulphonyl.

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Z particularly preferably represents the heterocyclic grouping below

 R^5

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particularly preferably represents hydrogen, hydroxyl, mercapto, cyano, fluorine, chlorine, bromine, iodine, represents in each case optionally fluorine-, chlorine-, methoxy-, ethoxy-, n- or i-propoxy-, n-, i-, s- or tbutoxy-, methylthio-, ethylthio-, n- or i-propylthio-, n-, i-, s- or t-butylthio-, methylsulphinyl-, ethylsulphinyl-, n- or i-propylsulphinyl-, methylsulphonyl-, ethylsulphonyl-, n- or i-propylsulphonyl-substituted methyl, ethyl, n- or ipropyl, n-, i-, s- or t-butyl, methoxy, ethoxy, n- or i-propoxy, n-, i-, s- or tbutoxy, methylthio, ethylthio, n- or i-propylthio, n-, i-, s- or t-butylthio, methylsulphinyl, ethylsulphinyl, n- or i-propylsulphinyl, methylsulphonyl, ethylsulphonyl, n- or i-propylsulphonyl, represents methylamino, ethylamino, n- or i-propylamino, n-, i-, s- or t-butylamino, dimethylamino, diethylamino, di-n-propylamino or di-i-propylamino, represents in each case optionally fluorine- and/or chlorine-substituted ethenyl, propenyl, butenenyl, ethinyl, propinyl, butinyl, propenyloxy, butenyloxy, propenylthio, butenylthio, propenylamino or butenylamino, represents in each case optionally fluorineand/or chlorine-substituted cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclopropyloxy, cyclobutyloxy, cyclopentyloxy, cyclohexyloxy, cyclopropylthio, cyclobutylthio, cyclopentylthio, cyclopentylthio, cyclopropylamino, cyclobutylamino, cyclopentylamino, cyclohexylamino, cyclopropylmethyl, cyclobutylmethyl, cyclopentylmethyl, cyclohexylmethyl, cyclopropylmethoxy, cyclobutylmethoxy, cyclopentylmethoxy, cyclohexylmethoxy, cyclopropylmethylthio, cyclobutylmethylthio, cyclopentylmethylthio, cyclohexylmethylthio, cyclopropylmethylamino, cyclobutylmethylamino, cyclopentylmethylamino or cyclohexylmethylamino, or represents in each case optionally fluorine-, chlorine-, methyl-, ethyl-, n- or i-propyl-, n-, i-, s- or t-butyl-, methoxy-, ethoxy-, n- or i-propoxy-substituted phenyl, phenyloxy, phenylthio, phenylamino, benzyl, benzyloxy, benzylthio or benzylamino, or in the case that two adjacent radicals R⁵ and R⁵ are located at a double bond together with the adjacent radical R⁵ also represents a benzo grouping,

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R6 particularly preferably represents hydrogen, hydroxyl, amino, represents in each case optionally fluorine- and/or chlorine-, methoxy- or ethoxysubstituted methyl, ethyl, n- or i-propyl, n-, i- or s-butyl, methoxy, ethoxy, nor i-propoxy, methylamino, ethylamino or dimethylamino, represents in each case optionally fluorine- and/or chlorine-substituted ethenyl, propenyl, ethinyl, propinyl or propenyloxy, represents in each case optionally fluorineand/or chlorine-substituted cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclopropylmethyl, cyclobutylmethyl, cyclopentylmethyl or cyclohexylmethyl, or represents in each case optionally fluorine-, chlorine-, methyl-, ethyl-, n- or i-propyl-, n-, i-, s- or t-butyl-, methoxy-, ethoxy-, n- or ipropoxy-substituted phenyl or benzyl, or together with an adjacent radical R⁵ or R6 represents in each case optionally methyl- and/or ethyl-substituted (trimethylene), butane-1,4-diyl (tetramethylene) propane-1,3-diyl pentane-1,5-diyl (pentamethylene),

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where the individual radicals R⁵ and R⁶ - if two or more of them are attached to the same heterocyclic groupings, may have identical or different meanings in the context of the above definition.

- 20 A very particularly preferably represents a single bond or represents methylene.
 - R1 very particularly preferably represents hydrogen, methyl, ethyl, n- or i-propyl.
 - R² very particularly preferably represents methyl.

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very particularly preferably represents hydrogen, nitro, cyano, fluorine, chlorine, bromine, iodine, methyl, ethyl, trifluoromethyl, methoxymethyl, methylthiomethyl, methylsulphinylmethyl, methylsulphonylmethyl, methoxy, ethoxy, difluoromethoxy, trifluoromethoxy, methylthio, ethylthio, methylsulphinyl, ethylsulphinyl, ethylsulphonyl, ethylsulphonyl or dimethylaminosulphonyl.

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 R^5

R4 very particularly preferably represents nitro, cyano, fluorine, chlorine, bromine, methyl, ethyl, trifluoromethyl, methoxymethyl, methylthiomethyl, methylsulphinylmethyl, methylsulphonylmethyl, methoxy, ethoxy, difluoromethoxy, trifluoromethoxy, methylthio, ethylthio, methylsulphinyl, ethylsulphonyl, ethylsulphonyl or dimethylaminosulphonyl.

very particularly preferably represents hydrogen, hydroxyl, chlorine, bromine, methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, difluoromethyl, dichloromethyl, trifluoromethyl, trichloromethyl, chlorodifluoromethyl, fluorodichloromethyl, fluoroethyl, chloroethyl, difluoroethyl, dichloroethyl, fluoron-propyl, fluoro-i-propyl, chloro-i-propyl, methoxymethyl, ethoxymethyl, methoxyethyl, methoxy, ethoxy, n- or i-propoxy, n-, i-, s- or t-butoxy, fluoroethoxy, chloroethoxy, difluoroethoxy, dichloroethoxy, trifluoroethoxy, trichloroethoxy, chlorofluoroethoxy, chlorodifluoroethoxy, fluorodichloroethoxy, methylthio, ethylthio, n- or i-propylthio, fluoroethylthio, chloroethylthio, difluoroethylthio, dichloroethylthio, methylsulphinyl, ethylsulphinyl, n- or i-propylsulphinyl, methylsulphonyl, ethylsulphonyl, dimethylamino, propenylthio, butenylthio, propinylthio, butinylthio, cyclopropyl, cyclopropylmethyl, cyclopropylmethoxy, phenyl or phenoxy.

very particularly preferably represents amino, methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, methoxy, ethoxy, methylamino, dimethylamino, cycloproypyl or cyclopropylmethyl, or together with R⁵ represents propane-1,3-diyl (trimethylene), butane-1,4-diyl (tetramethylene) or pentane-1,5-diyl (pentamethylene).

30 A most preferably represents methylene.

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The invention preferably provides the sodium, potassium, magnesium, calcium, ammonium, C_1 - C_4 -alkyl-ammonium-, di- $(C_1$ - C_4 -alkyl)-ammonium-, tri- $(C_1$ - C_4 -alkyl)-ammonium-, tetra- $(C_1$ - C_4 -alkyl)-ammonium, tri- $(C_1$ - C_4 -alkyl)-sulphonium, C_5 - or C_6 -cycloalkyl-ammonium and di- $(C_1$ - C_2 -alkyl)-benzyl-ammonium salts of the compounds of the formula (I), in which m, n, A, R^1 , R^2 , R^3 , R^4 and Z are each as defined above.

Preference according to the invention is given to compounds of the formula (I) which contain a combination of the meanings mentioned above as being preferred.

Particular preference according to the invention is given to compounds of the formula (I) which contain a combination of the meanings listed above as being particularly preferred.

Very particular preference according to the invention is given to the compounds of the formula (I) which contain a combination of the meanings listed above as being very particularly preferred.

Compounds of the general formulae (IA), (IB) and (IC) below are particularly emphasized as being according to the invention:

$$(R^2)_m$$
 R^1
 $(R^4)_n$
 (IB)

$$(R^{4})_{n}$$

$$(R^{2})_{m}$$

$$R^{1}$$

$$Q$$

$$N$$

$$R^{3}$$

$$N$$

$$R^{5}$$
(IC)

in which

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m represents the numbers 0, 1 or 2,

n represents the numbers 0, 1 or 2,

A particularly preferably represents a single bond or represents methylene,

10 Q represents oxygen or sulphur,

R¹ represents hydrogen, methyl, ethyl, n- or i-propyl,

 R^2 represents methyl,

D3 rapraganta hydro

R³ represents hydrogen, nitro, cyano, fluorine, chlorine, bromine, iodine, methyl, ethyl, trifluoromethyl, methoxymethyl, methylthiomethyl, methyl-sulphin-ylmethyl, methylsulphonylmethyl, methoxy, ethoxy, difluoromethoxy, trifluoromethoxy, methylthio, ethylthio, methylsulphinyl, ethylsulphinyl, methylsulphonyl, ethylsulphonyl or dimethylaminosulphonyl.

R4 represents nitro, cyano, fluorine, chlorine, bromine, methyl, ethyl, trifluoromethyl, methoxymethyl, methylthiomethyl, methylsulphinylmethyl, methylsulphonylmethyl, methoxy, ethoxy, difluoromethoxy, trifluoromethoxy, methylthio, ethylthio, methylsulphinyl, ethylsulphinyl, methylsulphonyl, ethylsulphonyl or dimethylaminosulphonyl,

represents hydrogen, hydroxyl, chlorine, bromine, methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, difluoromethyl, dichloromethyl, trifluoromethyl, trichloromethyl, chlorodifluoromethyl, fluorodichloromethyl, fluoroethyl, chloroethyl, difluoroethyl, dichloroethyl, fluoro-n-propyl, fluoro-i-propyl, chloro-i-propyl, methoxymethyl, ethoxymethyl, methoxyethyl, methoxy, ethoxy, n- or i-propoxy, n-, i-, s- or t-butoxy, fluoroethoxy, chloroethoxy, difluoroethoxy, dichloroethoxy, trifluoroethoxy, trichloroethoxy, chlorofluoroethoxy, chlorodifluoroethoxy, fluorodichloroethoxy, methylthio, ethylthio, n- or i-propylthio, fluoroethylthio, chlorodifluoroethylthio, dichloroethylthio, chlorofluoroethylthio, chlorodichloroethylthio, fluorodichloroethylthio, methylsulphinyl, ethylsulphinyl, n- or i-propylsulphinyl, methylsulphonyl, ethylsulphonyl, n- or i-propylsulphonyl, dimethylamino, propenylthio, butenylthio, propinylthio, butinylthio, cyclopropyl, cyclopropylmethyl, cyclopropylmethoxy, phenyl or phenoxy, and

represents amino, methyl, ethyl, n- or i-propyl, n-, i-, s- or t-butyl, methoxy, ethoxy, methylamino, dimethylamino, cyclopropyl or cyclopropylmethyl, or together with R⁵ represents propane-1,3-diyl (trimethylene), butane-1,4-diyl (tetramethylene) or pentane-1,5-diyl (pentamethylene).

Here, very particular emphasis is given to the compounds of the formula (IA) in which A represents methylene.

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 R^5

The abovementioned general or preferred radical definitions apply both to the end products of the formula (I) and also, correspondingly, to the starting materials or intermediates required in each case for the preparation. These radical definitions can be combined with one another at will, i.e. including combinations between the given preferred ranges.

Examples of compounds of the general formula (I) according to the invention are listed in the groups below.

10 Group 1

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$$(IA-1)$$

$$(R^4)_n$$

$$R^5$$

Here, R^3 , $(R^4)_n$, R^5 and R^6 each have, for example, the meanings given in the table below:

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
H	-	CF ₃	CH ₃
F	-	CF ₃	CH ₃
Cl	-	CF ₃	CH ₃
Br	-	CF ₃	CH ₃
I	-	CF ₃	CH ₃
NO ₂	-	CF ₃	CH ₃
CN	-	CF ₃	CH ₃
CH ₃	-	CF ₃	CH ₃
OCH ₃	-	CF ₃	CH ₃

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
CF ₃	-	CF ₃	CH ₃
OCHF ₂	-	CF ₃	CH ₃
OCF ₃	-	CF ₃	CH ₃
SO ₂ CH ₃	-	CF ₃	CH ₃
H	-	OCH ₃	CH ₃
F	-	OCH ₃	CH ₃
Cl	-	OCH ₃	CH ₃
Br	-	OCH ₃	CH ₃
I	-	OCH ₃	CH ₃
NO ₂	-	OCH ₃	CH ₃
CN	-	OCH ₃	CH ₃
CH ₃	-	OCH ₃	CH ₃
OCH ₃	-	OCH ₃	CH ₃
CF ₃	-	OCH ₃	CH ₃
OCHF ₂	-	OCH ₃	CH ₃
OCF ₃	-	OCH ₃	CH ₃
SO ₂ CH ₃	-	OCH ₃	CH ₃
Н	-	SCH ₃	CH ₃
F	-	SCH ₃	CH ₃
Cl	-	SCH ₃	CH ₃
Br	-	SCH ₃	CH ₃
I	-	SCH,	CH ₃
NO ₂	-	SCH ₃	CH ₃
CN	-	SCH ₃	CH ₃
CH ₃	-	SCH ₃	CH ₃
OCH ₃	•	SCH ₃	CH ₃
CF ₃	-	SCH ₃	CH ₃
OCHF ₂	-	SCH ₃	CH ₃
OCF ₃	-	SCH ₃	CH ₃

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	-	SCH ₃	CH ₃
Н	-	OC ₂ H ₅	CH ₃
F	-	OC_2H_5	CH ₃
CI	-	OC_2H_5	CH ₃
Br	-	OC_2H_5	CH ₃
Ι	-	OC_2H_5	CH ₃
NO ₂	-	OC ₂ H ₅	CH ₃
CN	-	OC_2H_5	CH ₃
CH ₃	-	OC_2H_5	CH ₃
OCH ₃	-	OC ₂ H ₅	CH ₃
CF ₃	-	OC ₂ H ₅	CH ₃
OCHF ₂	-	OC_2H_5	CH ₃
OCF ₃	-	OC ₂ H ₅	CH ₃
SO ₂ CH ₃	-	OC ₂ H ₅	CH ₃
Н	-	$N(CH_3)_2$	CH ₃
F	-	N(CH ₃) ₂	CH ₃
Cl	-	$N(CH_3)_2$	CH ₃
Br	-	$N(CH_3)_2$	CH ₃
I	-	$N(CH_3)_2$	CH ₃
NO ₂	-	$N(CH_3)_2$	CH ₃
CN	-	$N(CH_3)_2$	CH ₃
CH,	-	$N(CH_3)_2$	CH ₃
OCH ₃	-	N(CH ₃) ₂	CH ₃
CF ₃	-	N(CH ₃) ₂	CH ₃
OCHF ₂	-	N(CH ₃) ₂	CH ₃
OCF ₃	_	$N(CH_3)_2$	CH ₃
SO ₂ CH ₃	-	$N(CH_3)_2$	CH ₃
Н	-	OCH ₃	

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
F	-	OCH ₃	
Cl	-	OCH ₃	
Br	-	OCH ₃	\triangle
I	-	OCH ₃	\triangle
NO_2	-	OCH ₃	\triangle
CN	-	OCH ₃	\triangle
CH ₃	-	OCH ₃	
OCH ₃	-	OCH ₃	\triangle
CF ₃	-	OCH ₃	
OCHF ₂	-	OCH ₃	
OCF ₃	-	OCH ₃	
SO ₂ CH ₃	-	OCH ₃	\triangle
Н	(3-) Cl	CF ₃	CH ₃
F	(3-) Cl	CH ₃	CH ₃
Cl	(3-) Cl	OCH ₃	CH ₃
Br	(3-) Cl	Br	\triangle

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(3-) Cl	CF ₃	CH ₃
NO ₂	(3-) Cl	CH ₃	CH ₃
Cl	(3-) Cl	SCH ₃	CH ₃
CH ₃	(3-) Cl	Cl	CH ₃
OCH ₃	(3-) Cl	OCH ₃	CH ₃
CF ₃	(3-) Cl	CF ₃	CH ₃
OCHF ₂	(3-) C1	CH ₃	CH ₃
OCF ₃	(3-) C1	CH ₃	CH ₃
SO ₂ CH ₃	(3-) Cl	OCH ₃	CH ₃

Group 2

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Here R^3 , $(R^4)_n$, R^5 and R^6 each have, for example, the meanings given in the table below:

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R^6
Cl	(2-) Cl	CF ₃	CH ₃
Cl	(2-) Cl	SCH_3	CH ₃
Cl	(2-) Cl	SC ₂ H ₅	CH ₃
Cl	(2-) CI	SC ₃ H ₇	CH ₃
Cl	(2-) Cl	SC ₃ H ₇ -i	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) Cl		CH ₃
		5	CII
Cl	(2-) CI		CH_3
		S	
Cl	(2-) Cl	LMCH3	CH ₃
		s	
Cl	(2-) Cl		CH ₃
		S	
Cl	(2-) Cl	$\overline{\nabla}$	CH ₃
		Y	
		s	
Cl	(2-) Cl	SCH=C=CH ₂	CH ₃
Cl	(2-) C1	SCH ₂ CN	CH ₃
Cl	(2-) Cl	SCH ₂ CH ₂ CN	CH ₃
C1	(2-) Cl	OCH ₃	CH ₃
Cl	(2-) Cl	OC ₂ H ₅	CH ₃
Cl	(2-) C1	OC ₃ H ₇	CH ₃
Cl	(2-) Cl	OC ₂ H ₇ -i	CH ₃
Cl	(2-) C1	OC ₄ H ₉	CH ₃
Cl	(2-) Cl	OCH ₂ CF ₃	CH ₃
Cl	(2-) Cl	∇	CH ₃
		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
		0	
Cl	(2-) Cl	OC ₆ H ₅	CH ₃
Cl	(2-) C1	Н	CH ₃

R³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) Cl	CH ₃	CH ₃
Cl	(2-) Cl	C_2H_5	CH ₃
Cl	(2-) Cl	C_3H_7	CH ₃
Cl	(2-) Cl	C ₃ H ₇ -i	CH ₃
Cl	(2-) Cl	C_4H_9	CH ₃
Cl	(2-) C1	C ₄ H ₉ -i	CH ₃
Cl	(2-) Cl	C ₄ H ₉ -s	CH ₃
Cl	(2-) Cl	C ₄ H ₉ -t	CH ₃
Cl	(2-) CI	\triangle	CH ₃
Cl	(2-) Cl	\triangle	CH ₃
Cl	(2-) Cl	CH=CHCH ₃	CH ₃
Cl	(2-) C1		CH ₃
Cl	(2-) C1	CI	CH ₃
Cl	(2-) C1		CH ₃
Cl	(2-) C1	$N(CH_3)_2$	CH,
Cl	(2-) C1	N	CH ₃
Cl	(2-) Cl	Cl	CH ₃
Cl	(2-) Cl	Br	CH ₃
SO ₂ CH ₃	(2-) Cl	CF ₃	CH ₃
SO ₂ CH ₃	(2-) Cl	SCH ₃	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ^s	R ⁶
SO ₂ CH ₃	(2-) Cl	SC ₂ H ₅	CH ₃
SO ₂ CH ₃	(2-) C1	SC ₃ H ₇	CH ₃
SO ₂ CH ₃	(2-) Cl	SC ₃ H ₇ -i	CH ₃
SO ₂ CH ₃	(2-) C1		CH ₃
		S	
SO ₂ CH ₃	(2-) Cl		CH ₃
		S	
SO ₂ CH ₃	(2-) CI	LMCH³	CH ₃
		S	
SO ₂ CH ₃	(2-) Cl		CH ₃
		S	
SO ₂ CH ₃	(2-) Cl	∇	CH ₃
		s	
SO ₂ CH ₃	(2-) Cl	SCH=C=CH ₂	CH ₃
SO ₂ CH ₃	(2-) Cl	SCH ₂ CN	CH ₃
SO ₂ CH ₃	(2-) CI	SCH ₂ CH ₂ CN	CH ₃
SO ₂ CH ₃	(2-) Cl	OCH ₃	CH,
SO ₂ CH ₃	(2-) Cl	OC ₂ H ₅	CH ₃
SO ₂ CH ₃	(2-) Cl	OC ₃ H ₇	CH ₃
SO ₂ CH ₃	(2-) Cl	OC ₃ H ₇ -i	CH ₃
SO ₂ CH ₃	(2-) Cl	OC ₄ H ₉	CH ₃
SO ₂ CH ₃	(2-) Cl	OCH ₂ CF ₃	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	\mathbb{R}^6
SO ₂ CH ₃	(2-) Cl	∇	CH ₃
		0	
SO ₂ CH ₃	(2-) C1	OC ₆ H ₅	CH ₃
SO ₂ CH ₃	(2-) CI	H	CH ₃
SO ₂ CH ₃	(2-) Cl	CH ₃	CH ₃
SO ₂ CH ₃	(2-) Cl	C_2H_5	CH ₃
SO ₂ CH ₃	(2-) Cl	C ₃ H ₇	CH ₃
SO ₂ CH ₃	(2-) C1	C ₃ H ₇ -i	CH ₃
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉	CH ₃
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉ -i	CH ₃
SO ₂ CH ₃	(2-) C1	C ₄ H ₉ -s	CH ₃
SO ₂ CH ₃	(2-) CI	C ₄ H ₉ -t	CH ₃
SO ₂ CH ₃	(2-) Cl	\triangle	CH ₃
SO ₂ CH ₃	(2-) Cl	\triangle	CH ₃
SO ₂ CH ₃	(2-) Cl	CH=CHCH ₃	CH ₃
SO ₂ CH ₃	(2-) Cl		CH ₃
SO ₂ CH ₃	(2-) C1	CI	CH ₃
SO ₂ CH ₃	(2-) Cl		CH ₃
SO ₂ CH ₃	(2-) Cl	N(CH ₃) ₂	CH ₃
SO ₂ CH ₃	(2-) Cl		CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	\mathbf{R}^6
SO ₂ CH ₃	(2-) Cl	Cl	CH ₃
SO_2CH_3	(2-) Cl	Br	CH ₃
Cl	(2-) SO ₂ CH ₃	CF ₃	CH ₃
Cl	(2-) SO ₂ CH ₃	SCH ₃	CH ₃
Cl	(2-) SO ₂ CH ₃	SC ₂ H ₅	CH ₃
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇	CH ₃
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇ -i	CH ₃
Cl	(2-) SO ₂ CH ₃		CH ₃
-		S	
Cl	(2-) SO ₂ CH ₃		CH ₃
		s	
Cl	(2-) SO ₂ CH ₃	LMCH3	CH ₃
		s	
Cl	(2-) SO ₂ CH ₃		CH ₃
		s	
Cl	(2-) SO ₂ CH ₃	∇	CH ₃
		s	
Cl	(2-) SO ₂ CH ₃	SCH=C=CH ₂	CH ₃
Cl	(2-) SO ₂ CH ₃	SCH ₂ CN	CH ₃
Cl	(2-) SO ₂ CH ₃	SCH ₂ CH ₂ CN	CH ₃
Cl	(2-) SO ₂ CH ₃	OCH3	CH ₃
Cl	(2-) SO ₂ CH ₃	OC ₂ H ₅	CH ₃
Cl	(2-) SO ₂ CH ₃	OC ₃ H ₇	CH ₃
Cl	(2-) SO ₂ CH ₃	OC ₃ H ₇ -i	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	\mathbb{R}^6
Cl	(2-) SO ₂ CH ₃	OC ₄ H ₉	CH ₃
Cl	(2-) SO ₂ CH ₃	OCH ₂ CF ₃	CH ₃
Cl	(2-) SO ₂ CH ₃	∇	CH ₃
Cl	(2-) SO ₂ CH ₃	OC ₆ H ₅	CH ₃
Cl	(2-) SO ₂ CH ₃	Н	CH ₃
Cl	(2-) SO ₂ CH ₃	CH ₃	CH ₃
Cl	(2-) SO ₂ CH ₃	C_2H_5	CH ₃
Cl	(2-) SO ₂ CH ₃	C_3H_7	CH ₃
Cl	(2-) SO ₂ CH ₃	C ₃ H ₇ -i	CH ₃
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉	CH ₃
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -i	CH ₃
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -s	CH ₃
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -t	CH ₃
Cl	(2-) SO ₂ CH ₃	\triangle	CH ₃
Cl	(2-) SO ₂ CH ₃	\searrow	CH ₃
Cl	(2-) SO ₂ CH ₃	CH=CHCH ₃	CH ₃
Cl	(2-) SO ₂ CH ₃		CH ₃
Cl	(2-) SO ₂ CH ₃	CI	CH ₃
Cl	(2-) SO ₂ CH ₃		CH ₃
Cl	(2-) SO ₂ CH ₃	N(CH ₃) ₂	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) SO ₂ CH ₃	N	CH ₃
Cl	(2-) SO ₂ CH ₃	. Cl	CH ₃
Cl	(2-) SO ₂ CH ₃	Br	CH ₃
Cl	(2-) Cl	CF ₃	\triangle
Cl	(2-) Cl	SCH ₃	\triangle
Cl	(2-) Cl	SC₂H₅	\triangle
Cl	(2-) Cl	SC ₃ H ₇	\triangle
Cl	(2-) Cl	SC ₃ H ₇ -i	\triangle
Cl	(2-) Cl	s	\triangle
Cl	(2-) Cl	S	\triangle
Cl	(2-) C1	S CH ₃	\triangle
Cl	(2-) Cl	s	

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) Cl	S	\triangle
Cl	(2-) Cl	SCH=C=CH ₂	\triangle
Cl	(2-) Cl	SCH₂CN	\triangle
Cl	(2-) Cl	SCH ₂ CH ₂ CN	\triangle
Cl	(2-) Cl	OCH ₃	\triangle
Cl	(2-) Cl	OC ₂ H ₅	\triangle
Cl	(2-) Cl	OC ₃ H ₇	\triangle
Cl	(2-) Cl	OC₃H₁-i	\triangle
Cl	(2-) Cl	OC₄H ₉	\triangle
Cl	(2-) Cl	OCH ₂ CF ₃	\triangle
Cl	(2-) Cl		\triangle
Cl	(2-) Cl	OC ₆ H ₅	\triangle

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) C1	Н	\triangle
Cl	(2-) CI	CH ₃	\triangle
Cl	(2-) Cl	C ₂ H ₅	\triangle
Cl	(2-) Cl	C_3H_7	\triangle
Cl	(2-) C1	C ₃ H ₇ -i	\triangle
Cl	(2-) CI	C ₄ H ₉	\triangle
Cl	(2-) CI	C ₄ H ₉ -i	\triangle
Cl	(2-) Cl	C ₄ H ₉ -s	\triangle
Cl	(2-) Cl	C ₄ H ₉ -t	\triangle
Cl	(2-) C1	\triangle	\triangle
Cl	(2-) C1		\triangle
Cl	(2-) Cl	CH=CHCH ₃	\triangle
Cl	(2-) Cl		

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) C1	CI	
Cl	(2-) Cl		
Cl	(2-) Cl	N(CH ₃) ₂	\triangle
Cl	(2-) C1	N	\triangle
Cl	(2-) Cl	Cl	\triangle
Cl	(2-) Cl	Br	\triangle
SO ₂ CH ₃	(2-) Cl	CF ₃	\triangle
SO ₂ CH ₃	(2-) CI	SCH ₃	\triangle
SO ₂ CH ₃	(2-) CI	SC ₂ H ₅	\triangle
SO ₂ CH ₃	(2-) C1	SC ₃ H ₇	\triangle
SO ₂ CH ₃	(2-) Cl	SC ₃ H ₇ -i	\triangle
SO ₂ CH ₃	(2-) Cl	s	

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	(2-) Cl	S	\triangle
SO ₂ CH ₃	(2-) C1	S CH ₃	
SO ₂ CH ₃	(2-) Cl	s	
SO ₂ CH ₃	(2-) CI	S	
SO ₂ CH ₃	(2-) Cl	SCH=C=CH ₂	\triangle
SO ₂ CH ₃	(2-) Cl	SCH₂CN	\triangle
SO ₂ CH ₃	(2-) C1	SCH ₂ CH ₂ CN	\triangle
SO ₂ CH ₃	(2-) Cl	OCH ₃	\triangle
SO ₂ CH ₃	(2-) Cl	OC ₂ H ₅	
SO ₂ CH ₃	(2-) Cl	OC ₃ H ₇	\triangle
SO ₂ CH ₃	(2-) Cl	OC ₃ H ₇ -i	\triangle
SO ₂ CH ₃	(2-) Cl	OC₄H ₉	\triangle

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	(2-) Cl	OCH ₂ CF ₃	\triangle
SO ₂ CH ₃	(2-) Cl		
SO ₂ CH ₃	(2-) Cl	OC ₆ H ₅	
SO ₂ CH ₃	(2-) Cl	Н	\triangle
SO ₂ CH ₃	(2-) Cl	CH ₃	\triangle
SO ₂ CH ₃	(2-) Cl	C ₂ H ₅	\triangle
SO ₂ CH ₃	(2-) Cl	C_3H_7	\triangle
SO ₂ CH ₃	(2-) Cl	C ₃ H ₇ -i	\triangle
SO ₂ CH ₃	(2-) CI	C ₄ H ₉	\triangle
SO ₂ CH ₃	(2-) C1	C₄H₀-i	\triangle
SO ₂ CH ₃	(2-) Cl	C₄H ₉ -s	
SO ₂ CH ₃	(2-) Cl	C₄H ₉ -t	
SO ₂ CH ₃	(2-) Cl	\triangle	

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	(2-) Cl	\bigcirc	\triangle
SO ₂ CH ₃	(2-) Cl	CH=CHCH ₃	\triangle
SO ₂ CH ₃	(2-) CI		
SO ₂ CH ₃	(2-) CI	CI	
SO ₂ CH ₃	(2-) CI		
SO ₂ CH ₃	(2-) C1	N(CH ₃) ₂	\triangle
SO ₂ CH ₃	(2-) Cl		
SO ₂ CH ₃	(2-) Cl	Cl	\triangle
SO ₂ CH ₃	(2-) Cl	Br	\triangle
Cl	(2-) SO ₂ CH ₃	CF ₃	
Cl	(2-) SO ₂ CH ₃	SCH ₃	\triangle
Cl	(2-) SO ₂ CH ₃	SC ₂ H ₅	\triangle

R ³	(position-)(R ⁴) _n	R ⁵	\mathbb{R}^6
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇	\triangle
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇ -i	
Cl	(2-) SO ₂ CH ₃	s	\triangle
Cl	(2-) SO ₂ CH ₃	S	
Cl	(2-) SO ₂ CH ₃	S CH3	
CI	(2-) SO ₂ CH ₃	s	\triangle
Cl	(2-) SO ₂ CH ₃	s	
Cl	(2-) SO ₂ CH ₃	SCH=C=CH ₂	\triangle
Cl	(2-) SO ₂ CII ₃	SCH₂CN	
Cl	(2-) SO ₂ CH ₃	SCH ₂ CH ₂ CN	
CI	(2-) SO ₂ CH ₃	OCH ₃	

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) SO ₂ CH ₃	OC ₂ H ₅	\triangle
Cl	(2-) SO ₂ CH ₃	OC ₃ H ₇	\triangle
Cl	(2-) SO ₂ CH ₃	OC₃H₁-i	
CI	(2-) SO ₂ CH ₃	OC₄H ₉	\triangle
Cl	(2-) SO ₂ CH ₃	OCH ₂ CF ₃	\triangle
CI	(2-) SO ₂ CH ₃		
Cl	(2-) SO ₂ CH ₃	OC ₆ H ₅	\triangle
Cl	(2-) SO ₂ CH ₃	Н	\triangle
Cl	(2-) SO ₂ CH ₃	CH ₃	
CI	(2-) SO ₂ CH ₃	C ₂ H ₅	
Cl	(2-) SO ₂ CH ₃	C ₃ H ₇	\triangle
Cl	(2-) SO ₂ CH ₃	C ₃ H ₇ -i	\triangle
Cl	(2-) SO ₂ CH ₃	C₄H ₉	

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -i	\triangle
Cl	(2-) SO ₂ CH ₃	C₄H ₉ -s	\triangle
Cl	(2-) SO ₂ CH ₃	C₄H ₉ -t	\triangle
Cl	(2-) SO ₂ CH ₃	\triangle	\triangle
CI	(2-) SO ₂ CH ₃	\triangle	\triangle
Cl	(2-) SO ₂ CH ₃	CH=CHCH ₃	\triangle
Cl	(2-) SO ₂ CH ₃		\triangle
Cl	(2-) SO ₂ CH ₃	CI	\triangle
Cl	(2-) SO ₂ CH ₃		\triangle
Cl	(2-) SO ₂ CH ₃	N(CH ₃) ₂	\triangle
Cl	(2-) SO ₂ CH ₃	N	\triangle
Cl	(2-) SO ₂ CH ₃	Cl	\triangle

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) SO ₂ CH ₃	Br	\triangle
Cl	(2-) Cl	CF ₃	N(CH ₃) ₂
Cl	(2-) Cl	SCH ₃	N(CH ₃) ₂
Cl	(2-) Cl	SC ₂ H ₅	N(CH ₃) ₂
Cl	(2-) Cl	SC ₃ H ₇	N(CH ₃) ₂
Cl	(2-) Cl	SC ₃ H ₇ -i	N(CH ₃) ₂
Cl	(2-) Cl	s	N(CH ₃) ₂
Cl	(2-) Cl	s	N(CH ₃) ₂
Cl	(2-) Cl	S CH3	N(CH ₃) ₂
Cl	(2-) Cl	s	N(CH ₃) ₂
Cl	(2-) CI	s	N(CH ₃) ₂
Cl	(2-) Cl	SCH=C=CH ₂	N(CH ₃) ₂
Cl	(2-) Cl	SCH ₂ CN	N(CH ₃) ₂
Cl	(2-) Cl	SCH ₂ CH ₂ CN	N(CH ₃) ₂
Cl	(2-) Cl	OCH ₃	N(CH ₃) ₂
Cl	(2-) Cl	OC ₂ H ₅	N(CH ₃) ₂
Cl	(2-) Cl	OC ₃ H ₂	N(CH ₃) ₂
Cl	(2-) Cl	OC ₃ Hi	$N(CH_3)_2$

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) Cl	OC ₄ H ₉	$N(CH_3)_2$
CI	(2-) Cl	OCH ₂ CF ₃	$N(CH_3)_2$
Cl	(2-) C1		N(CH ₃) ₂
Cl	(2-) Cl	OC ₆ H ₅	N(CH ₃) ₂
Cl	(2-) Cl	Н	N(CH ₃) ₂
Cl	(2-) Cl	CH ₃	$N(CH_3)_2$
Cl	(2-) C1	C ₂ H ₅	N(CH ₃) ₂
Cl	(2-) C1	C_3H_7	N(CH ₃) ₂
Cl	(2-) Cl	C ₃ H ₇ -i	N(CH ₃) ₂
Cl	(2-) Cl	C ₄ H ₉	N(CH ₃) ₂
Cl	(2-) Cl	C ₄ H ₉ -i	N(CH ₃) ₂
Cl	(2-) Cl	C ₄ H ₉ -s	N(CH ₃) ₂
Cl	(2-) Cl	C ₄ H ₉ -t	N(CH ₃) ₂
Cl	(2-) Cl	\triangle	N(CH ₃) ₂
Cl	(2-) CI	\searrow	N(CH ₃) ₂
Cl	(2-) CI	CH=CHCH ₃	$N(CH_3)_2$
Cl	(2-) CI		N(CH ₃) ₂
Cl	(2-) Cl	CI	N(CH ₃) ₂
Cl	(2-) C1		N(CH ₃) ₂
Cl	(2-) Cl	N(CH ₃) ₂	N(CH ₃) ₂

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	\mathbb{R}^6	
Cl	(2-) C1	N	N(CH ₃) ₂	
Cl	(2-) Cl	Cl	$N(CH_3)_2$	
Cl	(2-) Cl	Br	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	CF ₃	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	SCH ₃	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	SC_2H_5	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	SC_3H_7	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	SC ₃ H ₇ -i	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) C1	s	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	s	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	S CH3	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) CI	s	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) C1	s	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	SCH=C=CH ₂	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) C1	SCH ₂ CN	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) C1	SCH ₂ CH ₂ CN	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	OCH ₃	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	OC ₂ H ₅	$N(CH_3)_2$	

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	(2-) Cl	OC_3H_7	$N(CH_3)_2$
SO ₂ CH ₃	(2-) Cl	OC ₃ H ₇ -i	$N(CH_3)_2$
SO ₂ CH ₃	(2-) C1	OC ₄ H ₉	$N(CH_3)_2$
SO ₂ CH ₃	(2-) C1	OCH ₂ CF ₃	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	∇	N(CH ₃) ₂
SO ₂ CH ₃	(2-) C1	OC ₆ H ₅	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	Н	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	CH ₃	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C_2H_5	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C_3H_7	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C ₃ H ₇ -i	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉ -i	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉ -s	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	C ₄ H ₉ -t	N(CH ₃) ₂
SO ₂ CH ₃	(2-) C1	\triangle	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	\triangle	N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	CH=CHCH ₃	$N(CH_3)_2$
SO ₂ CH ₃	(2-) Cl		N(CH ₃) ₂
SO ₂ CH ₃	(2-) Cl	CI	N(CH ₃) ₂

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶	
SO ₂ CH ₃	(2-) Cl		N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	N(CH ₃) ₂	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) C1	N	N(CH ₃) ₂	
SO ₂ CH ₃	(2-) Cl	CI	$N(CH_3)_2$	
SO ₂ CH ₃	(2-) Cl	Br	$N(CH_3)_2$	
Cl	(2-) SO ₂ CH ₃	CF ₃	$N(CH_3)_2$	
Cl	(2-) SO ₂ CH ₃	SCH ₃	$N(CH_3)_2$	
Cl	(2-) SO ₂ CH ₃	SC ₂ H ₅	N(CH ₃) ₂	
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇	$N(CH_3)_2$	
Cl	(2-) SO ₂ CH ₃	SC ₃ H ₇ -i	$N(CH_3)_2$	
Cl	(2-) SO ₂ CH ₃	s	N(CH ₃) ₂	
Cl	(2-) SO ₂ CH ₃	s	N(CH ₃) ₂	
Cl	(2-) SO ₂ CH ₃	S CH ₃	N(CH ₃) ₂	
Cl	(2-) SO ₂ CH ₃	S	N(CH ₃) ₂	
CI	(2-) SO ₂ CH ₃	S	N(CH ₃) ₂	
CI	(2-) SO ₂ CH ₃	SCH=C=CH ₂	N(CH ₃) ₂	

\mathbb{R}^3	(position-)(R ⁴) _n	(position-)(R ⁴) _n R ⁵	
Cl	(2-) SO ₂ CH ₃	SCH ₂ CN	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	SCH ₂ CH ₂ CN	N(CH ₃) ₂
C1	(2-) SO ₂ CH ₃	OCH ₃	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OC ₂ H ₅	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OC ₃ H ₇	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OC ₃ H ₇ -i	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OC₄H ₉	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OCH ₂ CF ₃	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	∇	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	OC ₆ H ₅	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	Н	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	CH ₃	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	C_2H_5	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	C_3H_7	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	C ₃ H ₇ -i	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	C_4H_9	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -i	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -s	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	C ₄ H ₉ -t	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	\triangle	N(CH ₃ I ₂
Cl	(2-) SO ₂ CH ₃		N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	СН=СНСН3	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃		N(CH ₃) ₂

R ³	(position-)(R ⁴) _n	R ⁵	R ⁶
Cl	(2-) SO ₂ CH ₃	CI	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃		N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	N(CH ₃) ₂	$N(CH_3)_2$
Cl	(2-) SO ₂ CH ₃	N	N(CH ₃) ₂
CI	(2-) SO ₂ CH ₃	Cl	N(CH ₃) ₂
Cl	(2-) SO ₂ CH ₃	Br	N(CH ₃) ₂
Cl	(2-) Cl	CH ₃	OCH ₃
Cl	(2-) Cl	C_2H_5	OCH ₃
Cl	(2-) Cl	C_3H_7	OCH ₃
Cl	(2-) Cl	SCH ₃	OCH ₃
Cl	(2-) Cl	SC ₂ H ₅	OCH ₃
Cl	(2-) CI	OCH ₃	OCH ₃
Cl	(2-) Cl	OC ₂ H ₅	OCH ₃
Cl	(2-) Cl	CH ₃	OC ₂ H ₅
Cl	(2-) Cl	C ₂ H ₅	OC ₂ H ₅
Cl	(2-) Cl	C_3H_7	OC ₂ H ₅
C1	(2-) CI	SCH ₃	OC_2H_5
Cl	(2-) Cl	SC ₂ H ₅	OC ₂ H ₅
Cl	(2-) Cl	OCH ₃	OC ₂ H ₅
Cl	(2-) Cl	OC ₂ H ₅	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	CH ₃	OCH ₃
Cl	(2-) SO ₂ CH ₃	C_2H_5	OCH ₃
Cl	(2-) SO ₂ CH ₃	C_3H_7	OCH ₃
Cl	(2-) SO ₂ CH ₃	SCH ₃	OCH ₃

\mathbb{R}^3	R ³ (position-)(R ⁴) _n		R ⁶
Cl	(2-) SO ₂ CH ₃	SC ₂ H ₅	OCH ₃
Cl	(2-) SO ₂ CH ₃	OCH ₃	OCH ₃
Cl	(2-) SO ₂ CH ₃	OC ₂ H ₅	OCH ₃
Cl	(2-) SO ₂ CH ₃	CH ₃	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	C ₂ H ₅	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	C ₃ H ₇	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	SCH ₃	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	SC ₂ H ₅	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	OCH ₃	OC ₂ H ₅
Cl	(2-) SO ₂ CH ₃	OC ₂ H ₅	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	Cl	OCH ₃
SO ₂ CH ₃	(2-) Cl	Br	OCH ₃
SO ₂ CH ₃	(2-) Cl	CH ₃	OCH ₃
SO ₂ CH ₃	(2-) Cl	C_2H_5	OCH ₃
SO ₂ CH ₃	(2-) Cl	C_3H_7	OCH ₃
SO ₂ CH ₃	(2-) Cl	SCH ₃	OCH ₃
SO ₂ CH ₃	(2-) Cl	SC ₂ H ₅	OCH ₃
SO ₂ CH ₃	(2-) C1	OCH ₃	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	OC ₂ H ₅	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	CH ₃	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	C_2H_5	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	C ₃ H ₇	OC ₂ H ₅
SO ₂ CH ₃	(2-) Cl	SCH ₃	OC ₂ H ₅
SO ₂ CH ₃	(2-) C1	SC ₂ H ₅	OC ₂ H ₅
SO ₂ CH ₃	(2-) CI	OCH ₃	OC ₂ H ₅
CF ₃	(2-) Cl	Br	CH ₃
CF ₃	(2-) CI	SCH ₃	CH ₃
CF ₃	(2-) CI	OCH ₃	CH ₃
CF ₃	(2-) Cl	N(CH ₃) ₂ CH ₃	

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
CF ₃	(2-) C1	CF ₃	CH ₃
CF ₃	(2-) NO ₂	Br	CH ₃
CF ₃	(2-) NO ₂	SCH ₃	CH ₃
CF ₃	(2-) NO ₂	OCH ₃	CH ₃
CF ₃	(2-) NO ₂	N(CH ₃) ₂	CH ₃
CF ₃	(2-) NO ₂	CF ₃	CH ₃
CF ₃	(2-) CH ₃	Br	CH ₃
CF ₃	(2-) CH ₃	SCH ₃	CH ₃
CF ₃	(2-) CH ₃	OCH ₃	CH ₃
CF ₃	(2-) CH ₃	N(CH ₃) ₂	CH ₃
CF ₃	(2-) CH ₃	CF ₃	CH ₃
CF ₃	(2-) OCH ₃	Br	CH ₃
CF ₃	(2-) OCH ₃	SCH ₃	CH ₃
CF ₃	(2-) OCH ₃	OCH ₃	CH ₃
CF ₃	(2-) OCH ₃	N(CH ₃) ₂	CH ₃
CF ₃	(2-) OCH ₃	CF ₃	CH ₃
SO ₂ CH ₃	(2-) NO ₂	Br	CH ₃
SO ₂ CH ₃	(2-) NO ₂	SCH ₃	CH ₃
SO ₂ CH ₃	(2-) NO ₂	OCH ₃	CH ₃
SO ₂ CH ₃	(2-) NO ₂	$N(CH_3)_2$	CH ₃
SO ₂ CH ₃	(2-) NO ₂	CF ₃	CH ₃
SO ₂ CH ₃	(2-) CF ₃	Br	CH ₃
SO ₂ CH ₃	(2-) CF ₃	SCH ₃	CH ₃
SO ₂ CH ₃	(2-) CF ₃	OCH ₃	CH ₃
SO ₂ CH ₃	(2-) CF ₃	N(CH ₃) ₂	CH ₃
SO ₂ CH ₃	(2-) CF ₃	CF ₃	CH ₃
SO ₂ CH ₃	(2-) SO ₂ CH ₃	Br	CH ₃
SO ₂ CH ₃	(2-) SO ₂ CH ₃	SCH ₃	CH ₃
SO ₂ CH ₃	(2-) SO ₂ CH ₃	OCH ₃	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶
SO ₂ CH ₃	(2-) SO ₂ CH ₃	$N(CH_3)_2$	CH ₃
SO ₂ CH ₃	(2-) SO ₂ CH ₃	CF ₃	CH ₃
CN	(2-) Cl	Br	CH ₃
CN	(2-) Cl	SCH ₃	CH ₃
CN	(2-) Cl	OCH ₃	CH ₃
CN	(2-) Cl	N(CH ₃) ₂	CH ₃
CN	(2-) Cl	CF ₃	CH ₃
CN	(2-) NO ₂	Br	CH ₃
CN	(2-) NO ₂	SCH ₃	CH ₃
CN	(2-) NO ₂	OCH ₃	CH ₃
CN	(2-) NO ₂	N(CH ₃) ₂	CH ₃
CN	(2-) NO ₂	CF ₃	CH ₃
CN	(2-) CF ₃	Br	CH ₃
CN	(2-) CF ₃	(2-) CF ₃ SCH ₃	
CN	(2-) CF ₃	OCH ₃	CH ₃
CN	(2-) CF ₃	N(CH ₃) ₂ CH	
CN	(2-) CF ₃	CF ₃	CH ₃
CN	(2-) SO ₂ CH ₃	Br	CH ₃
CN	(2-) SO ₂ CH ₃	SCH ₃	CH ₃
CN	(2-) SO ₂ CH ₃	OCH ₃	CH ₃
CN	(2-) SO ₂ CH ₃	N(CH ₃) ₂	CH ₃
CN	(2-) SO ₂ CH ₃	CF ₃	CH ₃
Br	(2-) NO ₂	Br	CH ₃
Br	(2-) NO ₂	SCH ₃	CH ₃
Br	(2-) NO ₂	OCH ₃	CH ₃
Br	(2-) NO ₂	$N(CH_3)_2$	CH ₃
Br	(2-) NO ₂	CF ₃ CH ₃	
Br	(2-) CF ₃	Br	CH ₃
Br	(2-) CF ₃	SCH ₃	CH ₃

\mathbb{R}^3	(position-)(R ⁴) _n	R ⁵	R ⁶	
Br	(2-) CF ₃	OCH ₃	CH ₃	
Br	(2-) CF ₃	$N(CH_3)_2$	CH ₃	
Br	(2-) CF ₃	CF ₃	CH ₃	
Br	(2-) SO ₂ CH ₃	Br	CH ₃	
Br	(2-) SO ₂ CH ₃	SCH ₃	CH ₃	
Br	(2-) SO ₂ CH ₃	OCH ₃	CH ₃	
Br	(2-) SO ₂ CH ₃	$N(CH_3)_2$	CH ₃	
Br	(2-) SO ₂ CH ₃	CF ₃	CH ₃	
Br	(2-) CH ₃	Br	CH ₃	
Br	(2-) CH ₃	SCH ₃	CH ₃	
Br	(2-) CH ₃	OCH ₃	CH ₃	
Br	(2-) CH ₃	$N(CH_3)_2$	CH ₃	
Br	(2-) CH ₃	CF ₃	CH ₃	

Group 3

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$$(R^4)_n = \begin{pmatrix} 0 & 0 & 0 \\ 1 & 2 & N & R^6 \\ R^3 & R^5 & R^5 \end{pmatrix}$$

Here, R^3 , $(R^4)_n$, R^5 and R^6 each have, for example, the meanings given above in Group 1.

Group 4

Here, R^3 , $(R^4)_n$, R^5 und R^6 each have, for example, the meanings given above in Group 2.

The novel substituted benzoylcyclohexanediones of the general formula (I) have strong and selective herbicidal activity.

The novel substituted benzoylcyclohexanediones of the general formula (I) are obtained when 1,3-cyclohexanedione or its derivatives of the general formula (II),

$$(R^2)_m$$
 (II)

in which,

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 m, R^1 and R^2 are each as defined above,

are reacted with substituted benzoic acids of the general formula (III).

$$A - Z$$
 (III)

in which

n, A, R³, R⁴ and Z are each as defined above.

5 in the presence of a dehydrating agent, if appropriate in the presence of one or more reaction auxiliaries and if appropriate in the presence of a diluent,

and, if appropriate, the compounds of the formula (I) obtained in this manner are subsequently subjected in a customary manner, within the scope of the definition of the substituents, to electrophilic or nucleophilic or oxidation or reduction reactions, or the compounds of the formula (I) are converted in a customary manner into salts.

The compounds of the formula (I) can be converted into other compounds of the formula (I) in accordance with the definition above using customary methods, for example by nucleophilic substitution (for example R^5 : $Cl \rightarrow OC_2H_5$, SCH_3) or by oxidation (for example R^5 : $CH_2SCH_3 \rightarrow CH_2S(O)CH_3$).

In principle, the compounds of the general formula (I) can also be synthesized as shown schematically below:

Reaction of 1,3-cyclohexanedione or its derivatives of the general formula (II) - above - with reactive derivatives of the substituted benzoic acids of the general formula (III) - above - in particular with the corresponding carbonyl chlorides. carboxylic anhydrides, carboxylic acid cyanides, methyl carboxylates or ethyl carboxylates - if appropriate in the presence of reaction auxiliaries, such as, for example, triethylamine (and, if appropriate, additionally zinc chloride), and, if appropriate, in the presence of a diluent, such as, for example, methylene chloride:

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$$(R^{2})_{m} \xrightarrow{\downarrow} O \qquad (R^{4})_{n}$$

$$R^{3} \qquad (I)$$

(Y for example CN, Cl)

In the reactions outlined above for preparing the compounds of the general formula (I), there is, in addition to the desired C-benzoylation at the cyclohexanedione, also an O-benzoylation - cf. equation below (cf. Synthesis 1978, 925-927; Tetrahedron Lett. 37 (1996), 1007-1009, WO-A-91/05469). However, the O-benzoyl compounds formed in this process are, under the reaction conditions of the process according to the invention, isomerized to the corresponding C-benzoyl compounds of the formula (I).

$$(R^{2})_{m}$$

$$R^{1}$$

$$(R^{2})_{m}$$

Using, for example, 1,3-cyclohexanedione and 2-(3-carboxy-5-fluorobenzyl)-5-ethyl-4-methoxy-2,4-dihydro-3H-1,2,4-triazol-3-one as starting materials, the course of the reaction in the process according to the invention can be outlined by the following equation:

The formula (II) provides a general definition of the cyclohexanediones to be used as starting materials in the process according to the invention for preparing compounds of the formula (I). In the formula (II), m, R¹ and R² each preferably have those meanings which have already been given above, in connection with the description of the compounds of the formula (I) according to the invention, as being preferred, as being particularly preferred, or as being very particularly preferred for m, R¹ and R².

The starting materials of the general formula (II) are known and/or can be prepared by processes known per se.

The formula (III) provides a general definition of the substituted benzoic acids further to be used as starting materials in the process according to the invention for preparing compounds of the formula (I). In the formula (III), n, A, R³, R⁴ and Z each preferably have those meanings which have already been given above, in connection with the description of the compounds of the formula (I) according to the invention, as being preferred, as being particularly preferred, as being very particularly preferred or as being most preferred for n, A, R³, R⁴ and Z.

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Except for 2-(5-carboxy-2,4-dichloro-phenyl)-4-difluoromethyl-5-methyl-2,4-dihydro-3H-1,2,4-triazol-3-one - alias 2,4-dichloro-5-(4-difluoromethyl-4,5-dihydro-3-methyl-5-oxo-1H-1,2,4-triazol-1-yl)-benzoic acid (CAS-Reg.-No. 90208-77-8) and 2-(5-carboxy-2,4-dichloro-phenyl)-4,5-dimethyl-2,4-dihydro-3H-1,2,4-triazol-3-one - alias 2,4-dichloro-5-(4,5-dihydro-3,4-dimethyl-5-oxo-1H-1,2,4-triazol-1-yl)-benzoic acid (CAS-Reg.-No. 90208-76-7) - the starting materials of the general formula (III) have hitherto not been disclosed in the literature. Except for 2-(5-carboxy-2,4-dichloro-phenyl)-4-difluoromethyl-5-methyl-2,4-dihydro-3H-1,2,4-triazol-3-one and

2-(5-carboxy-2,4-dichloro-phenyl)-4,5-dimethyl-2,4-dihydro-3H-1,2,4-triazol-3-one (cf. JP-A-58225070 - quoted in Chem. Abstracts 100:209881, JP-A-02015069 - quoted in Chem. Abstracts 113:23929), they also form, as novel compounds, part of the subject matter of the present application.

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The novel substituted benzoic acids of the general formula (III), are obtained when benzoic acid derivatives of the general formula (IV),

in which

n, A, R³ and R⁴ and Z are each as defined above, and

Y represents cyano, carbamoyl, halogenocarbamoyl or alkoxycarbonyl,

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are reacted with water, if appropriate in the presence of a hydrolysis auxiliary, such as, for example, sulphuric acid, at temperatures between 50°C and 120°C (cf. the Preparation Examples).

- The benzoic acid derivatives of the general formula (IV) required as precursors are known and/or can be prepared by processes known per se (cf. DE-A-3839480. DE-A-4239296, EP-A-597360, EP-A-609734, DE-A-4303676, EP-A-617026, DE-A-4405614, US-A-5378681).
- The novel substituted benzoic acids of the general formula (III) are also obtained when halogeno(alkyl)benzoic acids of the general formula (V).

HO
$$(R^4)_n$$

$$(V)$$

in which

 $n,\,A,\,R^3$ and R^4 are each as defined above and

X represents halogen (in particular fluorine, chlorine or bromine)

are reacted with compounds of the general formula (VI)

$$H_{Z}$$
 (VI)

in which

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Z is as defined above,

if appropriate in the presence of a reaction auxiliary, such as, for example, triethylamine or potassium carbonate, and if appropriate in the presence of a diluent, such as, for example, acetone, acetonitrile, N,N-dimethyl-formamide or N,N-dimethyl-acetamide, at temperatures between 50°C and 200°C (cf. the Preparation Examples).

possible, similarly to the methods described above, to react appropriate nitriles, amides and esters - in particular the methyl esters or the ethyl esters - with compounds of the general formula (VI). By subsequent hydrolysis according to customary methods, for example by reaction with aqueous-ethanolic potassium hydroxide solution, it is then possible to obtain the corresponding substituted benzoic

Instead of the halogeno(alkyl)benzoic acids of the general formula (V), it is also

acids.

The halogeno(alkyl)benzoic acids of the formula (V) - or corresponding nitriles or esters - required as precursors are known and/or can be prepared by processes known per se (cf. EP-A-90369, EP-A-93488, EP-A-399732, EP-A-480641, EP-A-609798, EP-A-763524, DE-A-2126720, WO-A-93/03722, WO-A-97/38977, US-A-3978127, US-A-4837333).

The compounds of the general formula (VI) further required as precursors are known and/or can be prepared by processes known per se.

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The process according to the invention for preparing the novel substituted benzoylcyclohexanediones of the general formula (I) is carried out using a dehydrating agent. Here, suitable dehydrating agents are the customary chemicals which are suitable for binding water.

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Examples of these are dicyclohexylcarbodiimide and carbonyl-bis-imidazole.

A particularly suitable dehydrating agent is dicyclohexylcarbodiimide.

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The process according to the invention for preparing novel substituted benzoylcyclohexanediones of the general formula (I) is, if appropriate, carried out using a reaction auxiliary.

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Examples of these are sodium cyanide, potassium cyanide, acetone cyanohydrin, 2-cyano-2-(trimethylsilyloxy)-propane and trimethylsilyl cyanide.

The particularly suitable further reaction auxiliary is trimethylsilyl cyanide.

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The process according to the invention for preparing the novel substituted benzoylcyclohexanediones of the general formula (I) is, if appropriate, carried out using a further reaction auxiliary. Suitable further reaction auxiliaries for the process

according to the invention are, in general, basic organic nitrogen compounds, such as, for example, trimethylamine, triethylamine, tripropylamine, tributylamine, ethyldiisopropylamine, N,N-dimethyl-cyclohexylamine, dicyclohexylamine, ethyldicyclohexylamine, N,N-dimethyl-aniline, N,N-dimethyl-benzylamine, pyridine, 2-methyl-, 3-methyl-, 4-methyl-, 2,4-dimethyl-, 2,6-dimethyl-, 3,4-dimethyl- and 3,5-dimethyl-pyridine, 5-ethyl-2-methyl-pyridine, 4-dimethylamino-pyridine, N-methylpiperidine, 1,4-diazabicyclo[2.2.2]-octane (DABCO), 1,5-diazabicyclo[4.3.0]-non-5-ene (DBN), or 1,8-diazabicyclo[5.4.0]-undec-7-ene (DBU).

Suitable diluents for carrying out the process according to the invention are, in particular, inert organic solvents. These include, in particular, aliphatic, alicyclic or aromatic, optionally halogenated hydrocarbons, such as, for example, benzine, benzene, toluene, xylene, chlorobenzene, dichlorobenzene, petroleum ether, hexane, cyclohexane, dichloromethane, chloroform, tetrachloromethane or 1,2-dichloroethane; ethers, such as diethyl ether, diisopropyl ether, dioxane, tetrahydrofuran, ethylene glycol dimethyl ether or ethylene glycol diethyl ether; ketones, such as acetone, butanone or methyl isobutyl ketone; nitriles, such as acetonitrile, propionitrile or butyronitrile; amides, such as N,N-dimethylformamide, N,N-dimethylacetamide, N-methyl-formanilide, N-methyl-pyrrolidone or hexamethylphosphoric triamide; esters such as methyl acetate or ethyl acetate, sulphoxides, such as dimethylsulphoxide.

When carrying out the process according to the invention, the reaction temperatures can be varied within a relatively wide range. In general, the process is carried out at temperatures between 0°C and 150°C, preferably between 10°C and 120°C.

The process according to the invention is generally carried out under atmospheric pressure. However, it is also possible to carry out the process according to the invention under elevated or reduced pressure - in general between 0.1 bar and 10 bar.

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For carrying out the process according to the invention, the starting materials are generally employed in approximately equimolar amounts. However, it is also possible to use a relatively large excess of one of the components. The reaction is generally carried out in a suitable diluent in the presence of a dehydrating agent, and the reaction mixture is generally stirred at the required temperature for several hours. Work-up is carried out by customary methods (cf. the Preparation Examples).

The active compounds according to the invention can be used as defoliants, desiccants, haulm killers and, especially, as weedkillers. By weeds in the broadest sense, there are to be understood all plants which grow in locations where they are not wanted. Whether the substances according to the invention act as total or selective herbicides depends essentially on the amount used.

The active compounds according to the invention can be used, for example, in connection with the following plants:

<u>Dicotyledonous weeds of the genera:</u> Sinapis, Lepidium, Galium, Stellaria, Matricaria, Anthemis, Galinsoga, Chenopodium, Urtica, Senecio, Amaranthus, Portulaca, Xanthium, Convolvulus, Ipomoea, Polygonum, Sesbania, Ambrosia, Cirsium, Carduus, Sonchus, Solanum, Rorippa, Rotala, Lindernia, Lamium, Veronica, Abutilon, Emex, Datura, Viola, Galeopsis, Papaver, Centaurea, Trifolium, Ranunculus, Taraxacum.

<u>Dicotyledonous crops of the genera:</u> Gossypium, Glycine, Beta, Daucus, Phaseolus, Pisum, Solanum, Linum, Ipomoea, Vicia, Nicotiana, Lycopersicon, Arachis, Brassica, Lactuca, Cucumis, Cucurbita.

Monocotyledonous weeds of the genera: Echinochloa, Setaria, Panicum, Digitaria, Phleum, Poa, Festuca, Eleusine, Brachiaria, Lolium, Bromus, Avena, Cyperus, Sorghum, Agropyron, Cynodon, Monochoria, Fimbristylis, Sagittaria, Eleocharis,

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Scirpus, Paspalum, Ischaemum, Sphenoclea, Dactyloctenium, Agrostis, Alopecurus. Apera.

Monocotyledonous crops of the genera: Oryza, Zea, Triticum, Hordeum, Avena, Secale, Sorghum, Panicum, Saccharum, Ananas, Asparagus, Allium.

However, the use of the active compounds according to the invention is in no way restricted to these genera, but also extends in the same manner to other plants.

The compounds are suitable, depending on the concentration, for the total control of weeds, for example on industrial terrain and railway tracks, and on paths and open spaces with or without tree plantings. Equally, the compounds can be employed for the control of weeds in perennial crops for example forests, decorative tree plantings, orchards, vineyards, citrus groves, nut orchards, banana plantations, coffee 15 plantations, tea plantations, rubber plantations, oil palm plantations, cocoa plantations, soft fruit plantings and hopfields, in lawns, turf and pasture land, and for the selective control of weeds in annual crops.

The compounds of the formula (I) according to the invention are particularly suitable for the selective control of monocotyledonous and dicotyledonous weeds in monocotyledonous crops, both pre-emergence and postemergence.

The active compounds can be converted into the customary formulations, such as solutions, emulsions, wettable powders, suspensions, powders, dusting agents, pastes, soluble powders, granules, suspo-emulsion concentrates, natural and synthetic materials impregnated with active compound, and very fine capsules in polymeric substances.

These formulations are produced in a known manner, for example by mixing the active compounds with extenders, that is liquid solvents and/or solid carriers. 5

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optionally with the use of surfactants, that is emulsifying agents and/or dispersing agents and/or foam-forming agents.

If the extender used is water, it is also possible to employ for example organic solvents as auxiliary solvents. Suitable liquid solvents are essentially the following: aromatics, such as xylene, toluene or alkylnaphthalenes, chlorinated aromatics and chlorinated aliphatic hydrocarbons, such as chlorobenzenes, chloroethylenes or methylene chloride, aliphatic hydrocarbons, such as cyclohexane or paraffins, for example petroleum fractions, mineral and vegetable oils, alcohols, such as butanol or glycol and also their ethers and esters, ketones, such as acetone, methyl ethyl ketone, methyl isobutyl ketone or cyclohexanone, strongly polar solvents, such as dimethylformamide and dimethyl sulphoxide, and also water.

Suitable solid carriers are: for example ammonium salts and ground natural minerals, such as kaolins, clays, talc, chalk, quartz, attapulgite, montmorillonite or diatomaceous earth, and ground synthetic minerals, such as finely divided silica, alumina and silicates; suitable solid carriers for granules are: for example crushed and fractionated natural rocks such as calcite, marble, pumice, sepiolite and dolomite, and also synthetic granules of inorganic and organic meals, and granules of organic material such as sawdust, coconut shells, maize cobs and tobacco stalks; suitable emulsifying and/or foam-forming agents are: for example nonionic and anionic emulsifiers, such as polyoxyethylene fatty acid esters, polyoxyethylene fatty alcohol ethers, for example alkylaryl polyglycol ethers, alkylsulphonates, alkyl sulphates, arylsulphonates as well as protein hydrolysates; suitable dispersing agents are: for example lignin-sulphite waste liquors and methylcellulose.

Tackifiers such as carboxymethylcellulose and natural and synthetic polymers in the form of powders, granules or latexes, such as gum arabic, polyvinyl alcohol and polyvinyl acetate, as well as natural phospholipids, such as cephalins and lecithins, and synthetic phospholipids, can be used in the formulations. Other possible additives are mineral and vegetable oils.

It is possible to use colorants such as inorganic pigments, for example iron oxide, titanium oxide and Prussian Blue, and organic dyes, such as alizarin dyes, azo dyes and metal phthalocyanine dyes, and trace nutrients such as salts of iron, manganese, boron, copper, cobalt, molybdenum and zinc.

The formulations in general contain between 0.1 and 95 per cent by weight of active compound, preferably between 0.5 and 90%.

For the control of weeds, the active compounds according to the invention, as such or in the form of their formulations, can also be used as mixtures with known herbicides, finished formulations or tank mixes being possible.

Possible components for the mixtures are known herbicides, for example

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acetochlor, acifluorfen(-sodium), aclonifen, alachlor, alloxydim(-sodium), ametryne, amidochlor, amidosulfuron, anilofos, asulam, atrazine, azafenidin, azimsulfuron, benazolin(-ethyl), benfuresate, bensulfuron(-methyl), bentazon, benzofenap, benzoylprop(-ethyl), bialaphos, bifenox, bispyribac(-sodium), bromobutide, bromofenoxim, bromoxynil, butachlor, butroxydim, butylate, cafenstrole, caloxydim, carbetamide, carfentrazone(-ethyl), chlomethoxyfen, chloramben, chloridazon, chlorimuron(-ethyl), chlornitrofen, chlorsulfuron, chlortoluron, cinidon(-ethyl), cinmethylin, cinosulfuron, clethodim, clodinafop(-propargyl), clomazone, clomeprop, clopyralid, clopyrasulfuron(-methyl), cloransulam(-methyl), cumyluron, cyanazine, cybutryne, cycloate, cyclosulfamuron, cycloxydim, cyhalofop(-butyl), 2,4-D, 2,4-DB, 2,4-DP, desmedipham, diallate, dicamba, diclofop(-methyl), diclosulam, diethatyl(-ethyl), difenzoquat, diflufenican, diflufenzopyr, dimefuron, dimepiperate, dimethachlor, dimethametryn, dimethenamid, dimexyflam, dinitramine, diphenamid, diquat, dithiopyr, diuron, dymron, epoprodan, EPTC, esprocarb, ethalfluralin, ethametsulfuron(-methyl), ethofumesate, ethoxyfen, ethoxysulfuron, etobenzanid, fenoxaprop(-P-ethyl), flamprop(-isopropyl), flamprop(-isopropyl-L).

flamprop(-methyl), flazasulfuron, fluazifop(-P-butyl), fluazolate, flucarbazone, flufenacet, flumetsulam, flumiclorac(-pentyl), flumioxazin, flumipropyn, flumetsulam, fluometuron, fluorochloridone, fluoroglycofen(-ethyl), flupoxam, flupropacil, flurpyrsulfuron(-methyl, -sodium), flurenol(-butyl), fluridone, fluroxypyr(-meptyl), flurprimidol, flurtamone, fluthiacet(-methyl), fluthiamide, fomesafen, glufosinate-(-ammonium), glyphosate(-isopropylammonium), halosafen, haloxyfop(-ethoxyethyl), haloxyfop(-P-methyl), hexazinone, imazamethabenz(-methyl), imazamethapyr, imazamox, imazapic, imazapyr, imazaquin, imazethapyr, imazosulfuron, iodosulfuron, ioxynil, isopropalin, isoproturon, isouron, isoxaben, isoxachlortole, isoxaflutole, isoxapyrifop, lactofen, lenacil, linuron, MCPA, MCPP, mefenacet, mesotrione, metamitron, metazachlor, methabenzthiazuron, metobenzuron, metobromuron, (alpha-)metolachlor, metosulam, metoxuron, metribuzin, metsulfuron(methyl), molinate, monolinuron, naproanilide, napropamide, neburon, nicosulfuron, norflurazon, orbencarb, oryzalin, oxadiargyl, oxadiazon, oxasulfuron, oxaziclomefone, oxyfluorfen, paraquat, pelargonic acid, pendimethalin, pentoxazone, phenmedipham, piperophos, pretilachlor, primisulfuron(-methyl), procarbazone, prometryn, propachlor, propanil, propaquizafop, propisochlor, propyzamide, prosulfocarb, prosulfuron, pyraflufen(-ethyl), pyrazolate, pyrazosulfuron(-ethyl), pyributicarb, pyridate, pyriminobac(-methyl), pyrazoxyfen, pyribenzoxim, pyrithiobac(-sodium), quinchlorac, quinmerac, quinoclamine, quizalofop(-P-ethyl), quizalofop(-P-tefuryl), rimsulfuron, sethoxydim, simazine, simetryn, sulcotrione, sulfentrazone, sulfometuron(-methyl), sulfosate, sulfosulfuron, tebutam, tebuthiuron. tepraloxydim, terbuthylazine, terbutryn, thenylchlor, thiafluamide, thiazopyr, thidiazimin, thifensulfuron(-methyl), thiobencarb, tiocarbazil, tralkoxydim, triallate, triasulfuron, tribenuron(-methyl), triclopyr, tridiphane, trifluralin and triflusulfuron.

Mixtures with other known active compounds, such as fungicides, insecticides, acaricides, nematicides, bird repellents, plant nutrients and agents which improve soil structure, are also possible.

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The active compounds can be used as such, in the form of their formulations or in the use forms prepared therefrom by further dilution, such as ready-to-use solutions, suspensions, emulsions, powders, pastes and granules. They are used in the customary manner, for example by watering, spraying, atomizing, scattering.

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The active compounds according to the invention can be applied either before or after emergence of the plants. They can also be incorporated into the soil before sowing.

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The amount of active compound used can vary within a relatively wide range. It depends essentially on the nature of the desired effect. In general, the amounts used are between 1 g and 10 kg of active compound per hectare of soil surface, preferably between 5 g and 5 kg per ha.

The preparation and use of the active compounds according to the invention can be seen from the following examples.

Preparation Examples:

Example 1

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

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1.2 g (3.48 mmol) of 5-ethoxy-4-methyl-2-(2-carboxy-5-trifluoromethyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one are suspended in 30 ml of acetonitrile and, at room temperature (approximately 20°C), admixed with 0.39 g (3.48 mmol) of 1,3-cyclohexanedione and 0.76 g (3.7 mmol) of dicyclohexylcarbodiimide (DCC). The reaction mixture is stirred at room temperature overnight (approximately 15 hours) and then admixed with 1.0 ml (7.0 mmol) of triethylamine and 0.10 ml (1.39 mmol) of trimethylsilyl cyanide. After 3 hours at room temperature, the mixture is stirred with 100 ml of 5% strength aqueous sodium carbonate solution, the dicyclohexylurea that separates out is filtered off with suction and the alkaline aqueous phase is repeatedly extracted with ethyl acetate. The aqueous phase is then adjusted to pH 2 using 35% strength hydrochloric acid and extracted repeatedly with methylene chloride. The methylene chloride phases are dried over sodium sulphate and concentrated.

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This gives 0.8 g (52% of theory) of 5-ethoxy-4-methyl-2-[2-(2,6-dioxo-cyclohexyl-carbonyl)-5-trifluoromethyl-benzyl]-2,4-dihydro-3H-1,2,4-triazol-3-one as an amorphous residue.

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logP (determined at pH=2): 2.70.

Example 2

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A solution of 1.5 g (7.2 mmol) of dicyclohexylcarbodiimide in 40 ml of acetonitrile is added to a suspension of 2.15 g (6.5 mmol) of 2-(4-carboxy-3-chloro-phenyl)-4-methyl-5-trifluoromethyl-2,4-dihydro-3H-1,2,4-triazol-3-one, 0.83 g (7.2 mmol) of 1,3-cyclohexanedione and 40 ml of acetonitrile, and the reaction mixture is stirred at 20°C for 16 hours. 1.3 g (13 mmol) of triethylamine and 0.26 g (2.6 mmol) of trimethylsilyl cyanide are then added, and the reaction mixture is stirred at 20°C for a further 4 hours. The mixture is then stirred with 180 ml of 2% strength aqueous sodium carbonate solution and filtered off with suction. The mother liquor is extracted with ethyl acetate. The aqueous phase is then acidified using 2N hydrochloric acid and extracted with methylene chloride. The organic phase is dried, concentrated under water pump vacuum and digested with diethyl ether/petroleum ether. The resulting crystalline product is isolated by filtration with suction.

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This gives 1.6 g (59% of theory) of 2-[4-(2,6-dioxocyclohexylcarbonyl)-3-chlorophenyl]-4-methyl-5-trifluoromethyl-2,4-dihydro-3H-1,2,4-triazol-3-one of melting point 182°C.

logP (determined at pH=2): 3.13.

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By the methods of Preparation Examples 1 and 2, and in accordance with the general description of the preparation processes according to the invention, it is also possible

to prepare, for example, the compounds of the formula (I) - or of the formulae (IA-3), (IB-2), (IC-2) or (ID) - listed in Tables 1 and 2 below.

$$(IA-3)$$

$$(R^4)_n$$

$$R^5$$

$$\begin{array}{c|c}
 & O \\
 & O \\$$

$$(R^4)_n$$
 N
 R^6
 $(IC-2)$

$$R^{2})_{m} \xrightarrow{4} A Z \qquad (ID)$$

<u>Table 1:</u> Examples of compounds of the formulae (IA-3), (IB-2), (IC-2)

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R ⁶	(formula) physical data
3	-	Н	Н	CF ₃	CH ₃	(IC-2)
						$\log P = 2.41a$
4	CH ₂	CF ₃	Н	Δ	Δ	(IA-3)
						$\log P = 2.41a$
5	CH ₂	SO ₂ CH ₃	Н	Δ	\triangle	(IB-2)
						m.p.: 153°C
6	CH ₂	SO ₂ CH ₃	Н	СН3	CH ₃	(IA-3)
						m.p.: 162°C
7	CH ₂	Cl	Н	CH ₃	СН3	(IB-2)
						$\log P = 1.50^{a}$
8	CH ₂	Cl	Н	CF ₃	СН3	(IB-2)
						$\log P = 2.44^{a}$
9	CH ₂	Cl	Н	\triangle	Δ	(IB-2)
						$\log P = 2.23b$
10	CH ₂	Br	Н	C ₂ H ₅	OC ₂ H ₅	(IA-3)
						$\log P = 2.68a$
11	CH ₂	F	Н	OC ₂ H ₅	CH ₃	(IA-3)
						$\log P = 1.73^{a}$
12	CH ₂	F	Н	SCH ₃	CH ₃	(IA-3)
						$\log P = 1.99a$
13	CH ₂	F	Н	SO ₂ CH ₃	СН3	(IA-3)
						$\log P = 1.83^{a}$
14	CH ₂	Br	Н	СН3	СН3	(IB-2)
						$\log P = 1.57a$
15	CH ₂	Br	Н	OC ₂ H ₅	СН3	(IB-2)
						m.p.: 132°C

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R6	(formula) physical data
16	CH ₂	Br	Н	\wedge		(IB-2) $logP = 2.31a$
17	CH ₂	Cl	Н	OC ₂ H ₅	\triangle	(IA-3) $logP = 3.03a$
18	CH ₂	Cl	Н	CF ₃	CH ₃	(IA-3) $logP = 2.75a$
19	CH ₂	Cl	Н	C ₂ H ₅	OC ₂ H ₅	(IA-3) $logP = 2.60a$
20	СН2	NO ₂	Н	SCH ₃	СН3	(IA-3) $logP = 2.04a$
21	CH ₂	CF3	Н	OC ₂ H ₅	\triangle	(IA-3) $logP = 3.02a$
22	CH ₂	CF ₃	Н	C ₂ H ₅	OC ₂ H ₅	(IA-3) $logP = 2.91a)$
23	CH ₂	CF ₃	Н	SCH ₃	СН3	(IA-3) $logP = 2.59a$
24	СН2	OCH ₃	Н	OC ₂ H ₅	СН3	(IA-3) $logP = 1.99a$
25	СН2	OCH ₃	Н	C ₂ H ₅	OC ₂ H ₅	(IA-3) $logP = 2.18a$
26	CH ₂	Br	Н	OC ₂ H ₅	СН3	(IA-3) $logP = 2.46a$
27	СН2	Br	Н	CF ₃	СН3	(IA-3) $logP = 2.85a$
28	CH ₂	Н	Н	CF ₃	CH ₃	(IA-3) $logP = 2.33a$
29	СН2	CF ₃	Н	OCH3	СН3	(IA-3) $logP = 2.35a$

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R6	(formula) physical data
30	CH ₂	F	Н	CF3	СН3	(IA-3)
						$\log P = 2.47^{a}$
31	CH ₂	F	Н	C ₂ H ₅	OC ₂ H ₅	(IA-3)
						$\log P = 2.28a$
32	CH ₂	F	Н	OCH ₃	CH ₃	(IA-3)
						$\log P = 1.76^{a}$
33	CH ₂	Н	Н	OC ₂ H ₅	CH ₃	(IA-3)
						$\log P = 1.93a$
34	CH ₂	Н	Н	OCH ₃	CH ₃	(IA-3)
						$\log P = 1.61a$
35	-	Н	(2) CF ₃	CF ₃	CH ₃	(IC-2)
						m.p.: 190°C
36	-	Н	Н	CF ₃	CH ₃	(IA-3)
						$logP = 2.48^{a}$
37	_	Cl	Н	CF ₃	CH ₃	(IA-3)
						$\log P = 2.83^{\text{ a}}$
38	-	Н	(2) Cl	CH ₃	CH ₃	(IC-2)
						m.p.: 196°C
39	CH ₂	Cl	(2) Cl	CF ₃	CH ₃	(IB-2)
						$\log P = 2.79^{a}$
40	-	Br	Н	CF_3	CH ₃	(IA-3)
						$\log P = 2.90^{a}$
41	CH ₂	Cl	(2) Cl	SCH_3	CH ₃	(IB-2)
						$\log P = 2.38^{a}$
42	CH ₂	Cl	(2) Cl	OC ₂ H ₅	CH ₃	(IB-2)
						$\log P = 2.48^{a}$
43	CH ₂	Cl	(2) Cl	$\overline{\triangle}$		(IB-2)
						$\log P = 2.62^{a}$

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R ⁶	(formula) physical data
44	CH ₂	Cl	(2) Cl	OCH ₃	CH ₃	(IB-2)
						$\log P = 2.14^{a}$
45	CH ₂	Cl	(2) Cl	OC ₃ H ₇ -i	CH ₃	(IB-2)
						$\log P = 2.79^{a}$
46	CH ₂	Cl	(2) Cl	OCH ₂ CF ₃	CH ₃	(IB-2)
						$\log P = 2.84^{a}$
47	CH ₂	Cl	(2) Cl	Br	CH ₃	(IB-2)
-						$\log P = 2.26^{a}$
48	CH ₂	Cl	(2) Cl	Н	CH ₃	(IB-2)
						$\log P = 1.69^{a}$
49	CH ₂	Cl	(2) Cl	\triangle	CH ₃	(IB-2)
						$\log P = 2.25^{a}$
50	CH ₂	Cl	(2) Cl	$N(CH_3)_2$	CH ₃	(IB-2)
						$\log P = 2.18^{a}$
51	CH ₂	Cl	(2) Cl	CH ₃	CH ₃	(IB-2)
						$\log P = 1.79^{a}$
52	CH ₂	Cl	(2) Cl	$R^5 + R^6$:	$(CH_2)_4$	(IB-2)
						$\log P = 1.98^{a}$
53	CH ₂	Cl	(2) Cl	OCH ₃	\triangle	(IB-2)
						$\log P = 2.45^{a}$
54	CH ₂	Cl	(2) Cl	OC ₂ H ₅	\triangle	(IB-2)
						$\log P = 2.79^{a}$
55	CH ₂	Cl	(2) Cl	OC ₃ H ₇ -i	Δ	(IB-2)
						$\log P = 3.14^{a}$
56	CH ₂	Cl	(2) Cl	OCH ₂ CF ₃	\triangle	(IB-2)
						$\log P = 3.18^{a}$
57	СН2	Cl	(2) Cl	SCH ₃		(IB-2)
						$\log P = 2.77^{a}$

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R6	(formula) physical data
58	CH ₂	Cl	(2) Cl	N(CH ₃) ₂	\triangle	(IB-2) $\log P = 2.49^{a}$
59	CH ₂	Cl	(2) Cl	CH ₃	\triangle	$(IB-2)$ $logP = 2.09^{a}$
60	CH ₂	Cl	(2) Cl	C_2H_5	OC ₂ H ₅	$(IB-2)$ $logP = 2.65^{a}$
61	CH ₂	CF ₃	Н	CF ₃	CH ₃	$(IA-3)$ $logP = 3.06^{a}$
62	СН2	Н	Н	C_2H_5	OC ₂ H ₅	$(IA-3)$ $logP = 2.10^{a}$
63	CH ₂	Н	Н	SCH ₃	CH ₃	$\log P = 1.85^{a}$
64	CH ₂	H	Н	\triangle	\triangle	$(IA-3)$ $logP = 2.09^{a}$
65	CH ₂	Cl	(5) Cl	CF ₃	CH ₃	(IA-3) $logP = 3.24^{a}$
66	CH ₂	Н	Н	SO ₂ CH ₃	CH ₃	$(IA-3)$ $logP = 1.71^{-a}$
67	CH ₂	SO ₂ CH ₃	Н	OC ₂ H ₅	CH ₃	$(IA-3)$ $logP = 1.64^{a}$
68	CH ₂	Br	Н	R ⁵ + R ⁵ :	(CH ₂) ₄	$(IA-3)$ $logP = 1.64^{a}$
69	CH ₂	Br	Н	OC ₃ H ₇ -n	CH ₃	$(IA-3)$ $logP = 2.82^{a}$
70	CH ₂	Br	Н	OC ₃ H ₇ -i	CH ₃	$(IA-3)$ $logP = 2.84^{a}$
71	CH ₂	CF ₃	Н	OC ₃ H ₇ -i	CH ₃	(IA-3) $logP = 3.05^{a}$

Ex. No.	A	R ³	(position) (R ⁴) _n	R5	R ⁶	(formula) physical
						data
72	CH ₂	CF ₃	Н	OC_3H_7 -n	CH ₃	(IA-3)
						$\log P = 3.06^{a}$
73	CH ₂	Br	Н	Br	CH ₃	(IA-3)
					,	$\log P = 2.33^{a}$
74	CH ₂	CF ₃	Н	OC_3H_7 -i		(IA-3)
						$\log P = 3.38^{a}$
75	CH ₂	CF ₃	Н	CH ₂ OCH ₃	\wedge	(IA-3)
						$\log P = 2.53^{a}$
76	CH ₂	CF ₃	Н	CH ₂ OCH ₃	CH ₃	(IA-3)
						$\log P = 2.26^{a}$
77	CH ₂	I	Н	CF ₃	CH ₃	(IA-3)
						$\log P = 2.98^{a}$
78	CH ₂	Br	Н	SCH ₃	CH ₃	(IA-3)
						$\log P = 2.36^{a}$
79	CH ₂	Cl	Н	SCH ₃	CH ₃	(IA-3)
						$\log P = 2.30^{a}$
80	CH ₂	CF ₃	Н	CH ₃	CH ₃	(IA-3)
						$\log P = 2.06^{a}$
81	CH ₂	CF ₃	Н	OC_2H_5	C_2H_5	(IA-3)
						$\log P = 3.01^{a}$
82	CH ₂	CF ₃	Н	$N(CH_3)_2$	CH ₃	(IA-3)
						$\log P = 2.40^{a}$
83	CH ₂	CF ₃	Н	Br	CH ₃	(IA-3)
				!		$\log P = 2.54^{a}$
84	CH ₂	Н	(3) CH ₃	OC_2H_5	CH ₃	(IA-3)
				_		$\log P = 2.21^{a}$
85	CH ₂	Br	Н	\triangle	\triangle	(IA-3)
						$\log P = 2.62^{a}$

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R ⁶	(formula) physical data
86	CH ₂	Br	Н		CH ₃	$(IA-3)$ $logP = 2.99^{a}$
87	CH ₂	CF ₃	Н	SC ₂ H ₅	CH ₃	(IA-3)
	_					$\log P = 2.94^{a}$
88	CH ₂	CF ₃	Н	SC ₃ H ₇ -i	CH ₃	$\log P = 2.63^{a}$
89	CH ₂	CF ₃	Н	$R^5 + R^6$:	(CH ₂) ₄	(IA-3)
90	CH ₂	CF ₃	Н	OCH ₃		$\log P = 2.25^{a}$ (IA-3)
	-	3		,		$\log P = 2.65^{a}$
91	CH ₂	CF ₃	Н	OCH ₂ CF ₃	CH ₃	$(IA-3)$ $logP = 3.06^{a}$
92	CH ₂	CN	Н	CF ₃	CH ₃	(IA-3)
93	CH ₂	F	Н	N(CH ₃) ₂	CH ₃	$\log P = 2.29^{a}$ (IA-3)
	0112	-		- 1 (3/2	13	$\log P = 1.81^{a}$
94	CH ₂	F	Н	OC ₃ H ₇ -n	CH ₃	$(IA-3)$ $logP = 2.44^{a}$
95	CH ₂	F	Н	CH ₂ OCH ₃	CH ₃	(IA-3)
96	CH ₂	F	Н	OCH ₃		$\log P = 1.69^{a}$
90	CII	1		00113		$\log P = 2.05^{a}$
97	CH ₂	F	Н	OC ₂ H ₅	\triangle	$(IA-3)$ $logP = 2.39^{a}$
98	СН2	I	Н	OC ₂ H ₅	CH ₃	$\frac{\log F - 2.39}{(\text{IA-3})}$
						$\log P = 2.59^{a}$

Ex. No.	A	R ³	(position) (R ⁴) _n	R ⁵	R6	(formula) physical data
99	CH ₂	OCH ₃	(2) NO ₂	OC_2H_5	CH ₃	(IC-2)
						$\log P = 2.24^{a}$
100	CH ₂	OCH ₃	(2) NO ₂	SCH ₃	CH ₃	(IC-2)
						$\log P = 2.18^{a}$

<u>Table 2</u>: Examples of compounds of the formula (ID)

Ex. No.	A	(position) R ¹	(position) (R ²) _m	(position) R ³	(position)	(position) Z	physical data
ID-1	CH ₂	H	H	(2) Cl	(R ⁴) _n (4) Cl		logP =
	~			36.2		N N	4.26 a)
						N	4.20
					1	O N	
						(3) CH ₃	
ID-2	CH ₂	(5) CH ₃	(5) CH ₃	(4) CF ₃	Н	Ö	logP =
	_					N CH ₃	2.61 a)
						N N 2 1 13	
						CH ₃	
						(2)	
ID-3	CH ₂	Н	Н	(4) CF ₃	Н	Q	logP =
						N CH ₃	2.24 a)
						N N 3 3	
		1			:	H C O	
						H ₃ C `O	
ID-4	CH ₂	Н	Н	(4) CF ₃	Н	Q	logP =
						CH₃	2.63 4)
						CH ₃	
						(2) CH ₃	
ID-5	CH ₂	Н	Н	Н	Н	Ö	logP =
							2.35 a)
						N _N N	
						(2)	

Ex. No.	A	(position) R ¹	(position) (R ²) _m	(position) R ³	(position) (R ⁴) _n	(position) Z	physical data
ID-6	CH ₂	Н	Н	(4) CF ₃	Н	$N = CF_3$	logP = 3.77 a)
ID-7	СН ₂	(5) CH ₃	(5) CH ₃	(4) CF ₃	Н	$(2) \qquad \qquad 0 $	logP = 3.27 a)
ID-8	CH ₂	(5) CH ₃	(5) CH ₃	(4) CF ₃	Н	$\begin{array}{c} O \\ N \\ \longrightarrow \\ N = \\ SCH_3 \end{array}$	logP = 3.18 a)
ID-9	CH ₂	Н	Н	(4) Br	Н	(2) N N	logP = 2.92 a)
ID-10	СН ₂	Н	Н	(4) Br	н	(2) O	logP = 3.04 ^{a)}
ID-11	CH ₂	(5) CH ₃	(5) CH ₃	(2) Cl	(4) Cl	$ \begin{array}{c} O \\ N \longrightarrow OCH_3 \end{array} $ OCH ₃	m.p.: 140°C logP = 2.72 31
ID-12	CH ₂	(5) CH ₃	(5) CH ₃	(2) Cl	(4) Cl	$ \begin{array}{c} $	m.p.: 103°C logP = 3.08 a)

Ex. No.	A	(position) R ¹	(position) (R ²) _m	(position) R ³	(position) (R ⁴) _n	(position) Z	physical data
ID-13	CH ₂	(5) CH ₃	(5) CH ₃	(2) Cl	(4) Cl	N CH ₃	m.p.: 118°C
						SCH ₃	$\log P = 2.98^{a}$
ID-14	CH ₂	(5) CH ₃	(5) CH ₃	(2) Cl	(4) Cl	N CH ₃	m.p.: 132°C
						(3) CH ₃	$logP = 2.32^{a}$
ID-15	CH ₂	(5) CH ₃	(5) CH ₃	(2) Cl	(4) CI	N CH ₃	m.p.: 170°C
						(3) Br	$\log P = 2.86^{a}$
ID-16	CH ₂	(4) CH ₃	(4) CH ₃	(2) CI	(4) CI	$ \begin{array}{c} O \\ N \\ O \\ O$	logP = 2.78 a)
ID-17	CH ₂	(4) CH ₃	(4) CH ₃	(2) Cl	(4) Cl	$ \begin{array}{c} $	logP = 3.15 a)
ID-18	CH ₂	(4) CH ₃	(4) CH ₃	(2) Cl	(4) CI	O N — CH ₃ SCH ₃	logP = 3.06 a)
ID-19	CH ₂	(4) CH ₃	(4) CH ₃	(2) Cl	(4) Cl	$ \begin{array}{c} $	logP = 2.38 a)

Ex. No.	A	(position)	(position) (R ²) _m	(position) R ³	(position)	(position)	physical data
ID-20	CH ₂	(4) CH ₃	(4) CH ₃	(2) Cl	(R ⁴) _n (4) Cl	0	logP =
						N CH ₃	2.94 a)
						N=(
						(3) Br	
ID-21	CH ₂	(5) C_3H_7 -i	Н	(2) Cl	(4) Cl	9	logP =
						N CH ₃	3.12 a)
				<u>}</u>		N=(OCH ₃	
						(3)	
ID-22	CH ₂	(5) C ₃ H ₇ -i	Н	(2) Cl	(4) Cl	0	$\log P = 3.49^{a}$
						N CH ₃	3.49
						OC ₂ H ₅	
ID-23	CH ₂	(5) C ₃ H ₇ -i	Н	(2) Cl	(4) Cl	(3)	logP =
110-23	Cn ₂	$(3) C_3 \Pi_7$ -1	П	(2) C1	(4) (1		3.39 a)
						N CH ₃	
						SCH ₃	
ID-24	CH ₂	(5) C ₃ H ₇ -i	Н	(2) Cl	(4) CI	0	logP =
		. , , , ,		, ,	, , , = -	N_CH ₃	2.70 a)
						CH ₃	
ID-25	CH ₂	(5) C ₃ H ₇ -i	Н	(2) Cl	(4) Cl	Q	logP =
						N CH ₃	3.28 a)
						N=(
						(3) Br	-
ID-26	CH ₂	(5) CH ₃	Н	(2) CI	(4) Cl	O II	
						N CH3	
						N=	
						(3) OCH ₃	
ID-27	CH ₂	(5) CH ₃	Н	(2) Cl	(4) Cl	0	
						N CH ₃	
						N=(OC ₂ H ₅	
						(3)	

Ex. No.	A	(position) R1	(position) (R ²) _m	(position) R ³	(position) (R ⁴) _n	(position) Z	physical data
ID-28	СН2	(5) CH ₃	H	(2) Cl	(4) Cl	N CH ₃	
		:		*		SCH ₃	
ID-29	CH ₂	(5) CH ₃	Н	(2) Cl	(4) Cl	$ \begin{array}{c} $	
ID-30	CH ₂	(5) CH ₃	Н	(2) Cl	(4) Cl	$ \begin{array}{c} $	

Starting materials of formula (III):

Example (III-1)

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4.5 g (15 mmol) of 2-(3-chloro-4-cyano-phenyl)-4-methyl-5-trifluoromethyl-2,4-dihydro-3H-1,2,4-triazol-3-one are taken up in 80 ml of 60% strength sulphuric acid. and the mixture is heated under reflux for 6 hours. After cooling to room temperature, the resulting crystalline produce is isolated by filtration with suction.

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This gives 4.5 g (91% of theory) of 2-(3-carboxy-4-chloro-phenyl)-4-methyl-5-trifluoromethyl-2,4-dihydro-3H-1,2,4-triazol-3-one of melting point 223°C.

Example (III-2)

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2 g (4.9 mmol) of 5-bromo-4-methyl-2-(2-ethoxycarbonyl-5-trifluoromethyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one (compare Example IV-1) are dissolved in 30 ml of 10% strength ethanolic potassium hydroxide solution and heated under reflux for 2 hours. The reaction mixture is concentrated under water pump vacuum, taken up in 20 ml of water and acidified with dilute hydrochloric acid. The solid that precipitates out is filtered and dried.

This gives 1.2 g (71% of theory) of 5-ethoxy-4-methyl-2-(2-carboxy-5-trifluoro-methyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one as a solid product. logP: 2.18a)

Example (III-3)

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13.4 g (35 mmol) 4-methyl-5-trifluoromethyl-2-(2,6-dichloro-3-methoxycarbonyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one are initially charged in 60 ml of 1,4-dioxane, and a solution of 1.54 g (38,5 mmol) of sodium hydroxide in 20 ml of water is slowly metered in at room temperature. The reaction mixture is stirred at 60°C for 150 minutes and subsequently concentrated under water pump vacuum. The residue is dissolved in 100 ml of water, and the pH of the solution is adjusted to 1 by addition of conc. hydrochloric acid. The resulting crystalline product is isolated by filtration with suction.

This gives 11.7 g (90% of theory) of 4-methyl-5-trifluoromethyl-2-(2,6-dichloro-3-carboxy-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one of melting point 207°C.

By the methods of Examples (III-1) and (III-3), it is also possible to prepare, for example, the compounds of the general formula (III) listed in Table 2 below.

$$HO \xrightarrow{1 \atop 2} \begin{pmatrix} (R^4)_n \\ A - Z \end{pmatrix}$$
 (III)

<u>Table 2</u>: Examples of compounds of the formula (III)

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-4	(4-) Cl	Н	$N \longrightarrow N \longrightarrow CH_3$ CH_3	$logP = 1.39^{a}$
III-5	(4-) SO ₂ CH ₃	Н	N=N	$logP = 1.47^{a}$
III-6	(4-) F	H	N N N N N N N N N N	$logP = 1.73^{a}$
III-7	(4-) CF ₃	H	N= Br	$logP = 1.65^{a}$
III-8	(4-) Br	Н	$(2-)$ $N \longrightarrow N$ $N \longrightarrow N$ $N (CH_3)_2$	$logP = 1.74^{a}$
III-9	(4-) CF ₃	Н	$ \begin{array}{c} $	$logP = 2.43^{a}$

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\rm n}$	-A-Z	
III-10	(4-) CF ₃	Н	$ \begin{array}{c c} & O \\ & N \\ & O \\$	$\log P = 2.12^{a}$
III-11	(4-) CF ₃	Н	O N— N— CH ₃	$logP = 1.61^{a}$
III-12	(4-) CF ₃	Н	(2-) N CH ₃ N(CH ₃) ₂	$logP = 1.93^{a}$
III-13	(4-) CF ₃	Н	N—CH ₃ Br	$logP = 2.01^{a}$
III-14	(4-) CF ₃	Н	(2-)	$logP = 1.77^{a}$
III-15	(3-) CH ₃	Н	O N CH_3 OC_2H_5	$logP = 1.70^{a}$

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	
III-16	(4-) SO ₂ CH ₃	Н	N CH ₃	$logP = 1.07^{a}$
			(2-) SCH ₃	1D - 2.25 a)
III-17	(4-) CF ₃	Н	$N \longrightarrow N \longrightarrow CH_3$ $N \longrightarrow CH_3$ SC_2H_5	$logP = 2.35^{a}$
III-18	(4-) CF ₃	Н	$ \begin{array}{c c} & & \\$	$logP = 2.63^{a}$
III-19	(4-) CF ₃	H	(2-) OCH ₃	$logP = 2.13^{a}$
III-20	(4-) CF ₃	Н	(2-) N N N	$logP = 1.82^{a}$
III-21	(4-) CF ₃	Н	OCH ₂ CF ₃	$logP = 2.48^{a}$

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	D 1 73 3)
III-22	(4-) CF ₃	H	N CH ₃	$logP = 1.73^{a}$
III-23	(4-) CF ₃	Н	(2-)	$logP = 3.11^{a}$
			$N = CF_3$	
III-24	(4-) F	Н	(2-) N CH ₃	$logP = 1.43^{a}$
			N=(N(CH ₃) ₂	$\log P = 1.97^{a}$
III-25	(4-) F	H	N N CH_3 OC_3H_7 - n	
III-26	(4-) F	Н	(2-) N CH ₃ CH ₂ OCH ₃	logP = 1.30 a)
III-27	(4-) F	Н	N N OCH ₃	$logP = 1.63^{a}$

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	
III-28	(4-) F	Н	$ \begin{array}{c} O \\ N = \\ OC_2H_5 \end{array} $	$logP = 1.93^{a}$
			(2-)	
III-29	(4-) CF ₃	Н	CH ₃	$logP = 1.78^{a}$
			(2-)	
III-30	(2-) Cl	(4-) Cl	0	m.p.: 230°C
			N CH ₃	$logP = 1.63^{a}$
			(3-) SCH ₃	
III-31	(2-) Cl	(4-) Cl	O	m.p.: 190°C
			N—CH ₃	$logP = 1.73^{a}$
			OC_2H_5	
III-32	(2-) Cl	(4-) C1	Q	m.p.: 210°C
			N=N	$logP = 1.87^{a}$
			(3-)	
III-33	(2-) Cl	(4-) Cl	0	m.p.: 210°C
			N CH ₃	$logP = 1.43^{a}$
			(3-) OCH ₃	

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-34	(2-) Cl	(4-) Cl	O	m.p.: 164°C
			N CH ₃	$logP = 2.01^{a}$
			N=(
			(3-j) OC ₃ H ₇ -i	
III-35	(2-) Cl	(4-) Cl	(3-)	m.p.: 168°C
			l n	$logP = 2.04^{a}$
			N CH ₃	
			OCH ₂ CF ₃	
III-36	(2-) Cl	(4-) Cl	O	m.p.: 218°C
			N CH ₃	$logP = 1.53^{a}$
			(3-) Br	
III-37	(2-) Cl	(4-) Cl	Q	m.p.: 259°C
			N CH ₃	$logP = 0.98^{a}$
			(3-) H	
III-38	(2-) Cl	(4-) Cl	0	m.p.: 210°C
			N CH ₃	$\log P = 1.56^{a}$
			N= (3-)	
III-39	(2-) Cl	(4-) Cl	(3-)	m.p.: 197°C
			Q	$\log P = 1.51^{a}$
			N—CH ₃	
			N(CH ₃) ₂	

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	
III-40	(2-) Cl	(4-) Cl	0	m.p.: 262°C
			N CH ₃	$logP = 1.11^{a}$
			N=(3-)	
			(3-)	
III-41	(2-) Cl	(4-) Cl	0	m.p.: 249°C
			N N	$logP = 1.30^{a}$
			(3-)	
III-42	(2-) Cl	(4-) Cl	0	m.p.: 200°C
			N N N	$logP = 1.71^{a}$
			N=(
		4	(3-) OCH ₃	
III-43	(2-) Cl	(4-) Cl	=0	m.p.: 189°C
			N N	$logP = 2.01^{a}$
			$N = \langle OC_2H_5 \rangle$	
			(3-)	
III-44	(2-) Cl	(4-) Cl	0 = <	m.p.: 178°C
			N N	$logP = 2.28^{a}$
			$N = \langle OC_3H_7 - i \rangle$	
			(3-)	
III-45	(2-) Cl	(4-) Cl	(3-)	m.p.: 161°C
				$logP = 2.31^{a}$
			NNN	
			OCH ₂ CF ₃	
			3323.3	

Ex.	(position-) R ³	(position-)	(position-) -A-Z	physical data
No.	(2-) Cl	$\frac{(\mathbf{R}^4)_{\mathbf{n}}}{(4-) \text{ Cl}}$	0	m.p.: 200°C
III-46	(2-) C1	(4-) 01	N N	$logP = 1.98^{a}$
			(3-) SCH ₃	20100
III-47	(2-) Cl	(4-) Cl	0 1	m.p.: 201°C
				$logP = 1.39^{a}$
			(3-) CH ₃	
III-48	(2-) Cl	(4-) Cl	(3-)	m.p.: 207°C
	·			$logP = 1.77^{a}$
			$N = \begin{pmatrix} N & N \\ N & N \end{pmatrix}_2$	
			14(0113/2	
III-49	(2-) Cl	(4-) Cl	(3-)	m.p.: 140°C
			0	$logP = 1.88^{a}$
			$N = N - OC_2H_5$	
ė			C ₂ H ₅	
III-50	(4-)	Н	Q	m.p.: 154°C
	OCH ₂ CHF ₂		N CH ₃	$logP = 2.14^{a}$
			(2-) CF ₃	
III-51	Н	Н	O	m.p.: 214°C
			N N	$logP = 1.87^{a}$
			(2-)	

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	
III-52	Н	Н		m.p.: 194°C
			(2-) O	$logP = 2.07^{a}$
III-53	Н	Н	Q	m.p.: 181°C
			N CI	$logP = 1.97^{a}$
			(2-)	25100
III-54	Н	Н	S	m.p.: 251°C
			(2-) NH	$logP = 1.14^{a}$
III-55	(2-) Cl	(4-) Cl	O	$logP = 1.38^{a}$
			(3-) N CH ₃	1 40 3
III-56	(2-) Cl	(4-) Cl	(3-) CH ₃	$logP = 1.48^{a}$
III-57	(2-) Cl	(4-) Cl	N SO_2	
III-58	(4-) Cl	H	O N——N——CH ₃ CF ₃	¹ H NMR (DMSO-D6, δ): 5.42 ppm.

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	¹H NMR
III-59	(4-) CF ₃	H		
			N CH ₃	(DMSO-D6, δ):
			N=	5.48 ppm.
			(2-) CH ₃	
			(2-)	¹H NMR
III-60	$(4-) CF_3$	Н		
			N CH3	(DMSO-D6, δ):
			N=	5.60 ppm.
			(2-) CF ₃	$LogP = 2.47^{a}$
III-61	(4-) CF ₃	Н	0	$logP = 2.33^{a}$
ļ 				
			(2-)	
				¹H NMR
III-62	(4-) SO ₂ CH ₃	Н	O	
			N CH ₃	(DMSO-D6, δ):
			N=<	5.14 ppm.
			(3-)	
111 (2	(4) SO CH	H	0	¹H NMR
III-63	(4-) SO ₂ CH ₃	11	N CH ₃	(DMSO-D6, δ):
			N N N 13	
			N	5.27 ppm.
			(2-) CH ₃	
III-64	(4-) Cl	Н	O	H NMR (CDCl ₃ .
			N CH ₃	δ): 5.12 ppm.
			CH ₃	
			(3-)	

Ex.	(position-)	(position-)	(position-)	physical data
No.	$\frac{R^3}{(4) C!}$	(R ⁴) _n H	-A-Z	'H NMR
III-65	(4-) Cl	п	Ĭ au	(DMSO-D6, δ):
			N CH ₃	· ·
			N=	5.20 ppm.
			(3-) CF ₃	
III-66	(4-) Cl	Н	0	¹H NMR
			N	(DMSO-D6, δ):
			N=	5.03 ppm.
			(2-)	
III-67	(4-) Br	H	(2-)	'H NMR
111 07	(1)21		Q	(DMSO-D6, δ):
			$N \longrightarrow N \longrightarrow OC_2H_5$	5.24 ppm.
			N=(
			C ₂ H ₅	
III-68	(4-) Br	Н	O	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=	5.39 ppm.
			(2-) CF ₃	
III-69	(4-) F	Н	Q	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=(5.19 ppm.
			$(2-)$ OC_2H_5	
III-70	(4-) F	Н	Q	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=(5.30 ppm.
			(2-) SCH ₃	

Ex.	(position-)	(position-)	(position-)	physical data
No.	R ³	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	
III-71	(4-) F	Н	(2-)	¹H NMR
			O II	(DMSO-D6, δ):
			N CH ₃	5.43 ppm.
			SO₂CH₃	
III-72	(4-) Br	Н	0	¹ H NMR, (CDCl ₃
			N CH3	δ):
			N=	5.10 ppm.
			(3-) CH ₃	
III-73	(4-) Br	Н	O II	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=(5.03 ppm.
			(3-) OC ₂ H ₅	
III-74	(4-) Br	Н	Q	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=	5.19 ppm.
			(3-) CF ₃	
III-75	(4-) Br	Н	0	'H NMR
			NNNN	(DMSO-D6, δ):
			N=	5.01 ppm.
			(2-)	
III-76	(4-) Cl	Н	O	'H NMR
			N	(DMSO-D6, δ):
) N=	5.14 ppm.
			(2-) OC ₂ H ₅	

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	
III-77	(4-) Cl	Н	(2-)	¹H NMR
			0	(DMSO-D6, δ):
			N OC ₂ H ₅	5.25 ppm.
			C_2H_5	
III-78	(4-) NO ₂	Н	0	¹H NMR
			\sim_{N}	(DMSO-D6, δ):
			N=(5.23 ppm.
			(2-) OC ₂ H ₅	
III-79	(4-) NO ₂	Н	O	¹H NMR
			N CH ₃	(DMSO-D6, δ):
			N=(5.37 ppm.
			(2-)	
III-80	(4-) CF ₃	Н	0	$logP = 2.46^{a}$
			N N	
			OC ₂ H ₅	
III-81	(4-) CF ₃	Н	(2-)	¹H NMR
			O.	(DMSO-D6, δ):
			N OC ₂ H ₅	5.31 ppm.
			$N = \langle C_2H_5 \rangle$	
III-82	(4-) CF ₃	Н	0	$logP = 2.08^{a}$
			N CH ₃	
			N= SCH ₃	
			(2-)	

Ex.	(position-)	(position-)	(position-)	physical data
No.	R^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	
III-83	(4-) OCH ₃	Н	0	¹ H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.38 ppm.
			\ \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	,
İ			N=	
			(2-) OC ₂ H ₅	
III-84	(4-) OCH ₃	Н	(2-)	¹ H NMR (CDCl ₃ ,
111-04	(4-) 00113		0	δ): 5.43 ppm.
			Ĭ	О). 5. 15 ррил.
			N OC ₂ H ₅	
			N=(
			C ₂ H ₅	
III-85	(4-) CF ₃	Н	(2-)	¹ H NMR (CDCl ₃ ,
111-05	(4-) C1 3		(-)	δ): 5.47 ppm.
			\parallel	<i>o)</i> : 5. 17 ppin.
			N CH ₃	
			N=	
			CH ₂ OCH ₃	
TIT OC	(4) D=	Н	0	$LogP = 1.44^{a}$
III-86	(4-) Br	n	Ĭ	
			N N	
			N=	
			(2-)	
III-87	(4-) Br	Н	Q	$LogP = 1.63^{a}$
			N N	
			N-	
			(2-)	
III-88	(4-) Br	Н	O II	$LogP = 2.27^{a}$
			N CH ₃	
			N=(
			OC ₃ H ₇ -	i
			(2-)	

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	
III-89	(4-) Br	Н	(2-)	$LogP = 2.31^{a}$
			0	
			Ĭ	
			N CH ₃	
			N=(
			`OC₃H ₇ -n	
				T D 1 02 3)
III-90	H	Н		$LogP = 1.82^{a}$
			N CH ₃	
			N N 33	
			Ì —⟨	
			CF ₃	
			(2-)	
III-91	(4-) Br	Н	0	¹ H NMR (CDCl ₃ ,
111-71	(4-) Di	11		
			N CH ₃	δ): 5.32 ppm.
			N=(
			\	
			OC_2H_5	
				TINDAD (CDC)
III-92	(4-) Br	Н		¹ H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.53 ppm.
			N N 3	11
			N=(
			CF ₃	
			(2-)	
III-93	(4-) F	Н	Q	¹ H NMR (CDCl ₃ ,
111-93	(4-)1			
			N N CH ₃	δ): 5.39 ppm.
		!	, OC H	
			OC_2H_5	
			1	H NMR (CDCl ₃ ,
III-94	(4-) F	Н		n NWR (CDCI3,
			N N CH ₃	δ): 5.57 ppm.
			N N 13	
			N=(
			CF ₃	
			(2-)	

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-95	(4-) F	H	(2-)	¹ H NMR (CDCl ₃ ,
			Q	δ): 5.44 ppm.
			$N \sim N \sim OC_2H_5$	
			N=(
			C ₂ H ₅	
III-96	(4-) F	Н	0	¹ H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.41 ppm.
			N=(
			(2-) OCH ₃	
III-97	H	Н	Q	¹ H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.34 ppm.
			N=(
			OC ₂ H ₅	
III-98	Н	Н	Q	¹H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.38 ppm.
			(2-) OCH ₃	
III-99	Н	Н	0	¹ H NMR (CDCl ₃ ,
111-99	*1			δ): 5.26 ppm.
), 0.20 pp
			(2-)	
III-100	Н	Н	0	¹ H NMR (CDCl ₃ ,
			N CH ₃	δ): 5.43 ppm.
			N=	
			(2-) SCH ₃	
	1			1

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{n}$	-A-Z	r . D 1 22 a)
III-101	Н	Н	(2-)	$LogP = 1.23^{a_j}$
			0	
			Ĭ	
			N CH ₃	
			/	
			N=	
			`SO ₂ CH ₃	
	(1 00 01			$logP = 1.14^{a}$
III-102	(4-) SO ₂ CH ₃	Н		logr - 1.14
			N CH ₃	
			/ N N	
			N=	
			OC ₂ H ₅	
			(2-)	
III-103	(4-) CF ₃	Н	Q	$logP = 2.45^{a}$
			N CH ₃	
			\ \n=\	
			OC ₃ H ₇ -i	
			(2-)	
TIT 104	(4.) CE	Н	(2-)	$logP = 2.48^{a}$
III-104	(4-) CF ₃	П	(2-)	1081 2.10
			Q.	
			N CH ₃	
			\ \w=\	
			OC ₃ H ₇ -n	
			3, 17	
III-105	(4-) Br	H	Ö	$logP = 1.85^{a}$
111-103	(1)			-
			N CH ₃	
			\	
			Br	
			(2-)	
TIT 106	(A) CE	H	0	$logP = 2.74^{a}$
III-106	(4-) CF ₃	П	Ŭ ∧	1061 2
			\sim N N	
			N	
			OC_3H_7 -i	

Ex.	(position-)	(position-)	(position-)	physical data
No.	R ³	(R ⁴) _n	-A-Z	1 D 2 01 a)
III-107	(4-) CF ₃	Н	(2-)	$logP = 2.01^{a}$
			Q	
			N N	
) N=(
			CH ₂ OCH ₃	
			3.12-31.3	
III-108	(4-) CF ₃	Н	(2-)	$logP = 1.79^{a}$
	* * * * * * * * * * * * * * * * * *			
			l II	
			N CH ₃	
			\\\\\\	
			N—	
			,CH ⁵ OCH³	
III-109	(4-) CF ₃	Н	0	$logP = 1.65^{a}$
111-107	(4-) CI 3	11	Ŭ	1051
			N N CH ₃	
			\ \\=\	
			Br	
			(2-)	
III-110	(4-) Br	Н	Q	$logP = 1.90^{a}$
	, , ,		│	
			N N CH ₃	
			$N=\langle$	
			SCH ₃	
			(2-)	
III-111	(4-) Cl	Н	0	$logP = 1.83^{a}$
1112111			ĺ	
			N CH ₃	
			\ \ \ =⟨	
			SCH ₃	
			(2-)	
III-112	(4-) I	Н	0	$logP = 2.06^{a}$
111-112	(4-)1	11		1051 2.00
			N CH ₃	
			N=(
	1		OC-H-	
			(2-)	

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-113	(4-) I	H	(2-)	
			O.	
			N CH ₃	
			H ₃ C N=	
			C ₂ H ₅	
III-114	(4-) Br	Н	0	m.p.: 191°C
			N	
			(2-) N	
III-115	(4-) Br	Н	0	m.p.: 213°C
			N N	
			N _N	
		TT	(2-) N	
III-116	Н	Н		
			-N	
			(2-)	
TTT 117	Н	Н	0	m.p.: 112°C
III-117	n i	11	N CH ₃	m.p.: 112 C
			N N N N N N N N N N N N N N N N N N N	
			CF ₃	
			(2-)	
III-118	(4-) CF ₃	Н	0	m.p.: 158°C
			N N CH ₃	
			CF ₃	
			(2-)	
III-119	$(4-) CF_3$	Н		m.p.: 162°C
			N N	
			N=	
			(2-)	
			(2)	

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-120	(4-) Cl	(5-) Cl	Q Q	m.p.: 167°C
111 120	(1) 01		N N CH_3 CF_3	
			(2-)	
III-121	Н	Н	N—CH ₃	m.p.: 188°C
III-122	Н	Н	(2-) N	
III-123	Н	Н	N CH ₃	m.p.: 131°C
III-124	(4-) Cl	H	O N—CH ₃ CF ₃	m.p.: 109°C
III-125	(4-) I	Н	N N N CH_3 CF_3	m.p.: 104°C
III-126	(4-) Br	Н	N CH ₃ CF ₃	m.p.: 99°C

Ex. No.	(position-) R ³	(position-) (R ⁴) _n	(position-) -A-Z	physical data
III-127	(4-) Br	Н	N N N N N N N N N N N N N N N N N N N	m.p.: 174°C
III-128	Н	Н	N—N—CH ₃ SCH ₃	m.p.: 122°C
III-129	(4-) Br	Н	N_CH ₃ SCH ₃	m.p.: 164°C
III-130	Н	Н	N N N N N N N N N N	m.p.: 154°C
III-131	(4-) Br	Н	$ \begin{array}{c c} & O \\ & N - CH_3 \\ & OC_3H_7-i \end{array} $	m.p.: 161°C
III-132	(4-) CN	Н	O N— N— CF ₃	m.p.: 196°C
III-133	Н	Н	(2-) N	m.p.: 192°C

Ex.	(position-)	(position-)	(position-)	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	
III-134	Н	Н	0=N (2-)	
III-135	(4-) Br	Н	(2)	m.p.: 252°C
III-136	(2-) NO ₂	(3-) OCH ₃	$ \begin{array}{c c} O \\ N \\ \hline OC_2H_5 \end{array} $	$logP = 1.65^{a}$
III-137	(2-) NO ₂	(3-) OCH ₃	N N N N N N N N N N	$\log P = 1.58^{a}$

Starting materials of the formula (IV):

Example (IV-1)

$$O \longrightarrow OC_2H_5 O \longrightarrow N \longrightarrow CH_3$$

$$CF_3$$

$$Br$$

Step 1

5

10 g (49 mmol) of 2-methyl-4-trifluoromethyl-benzoic acid are dissolved in 150 ml of ethanol and admixed with 1 ml of conc. sulphuric acid. The solution is heated under reflux for 24 hours and then concentrated, and the residue is taken up in methylene chloride and extracted with saturated aqueous sodium bicarbonate solution. The methylene chloride phase is dried over sodium sulphate and concentrated under water pump vacuum.

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This gives 9 g (80% of theory) of ethyl 2-methyl-4-trifluoromethyl-benzoate as an amorphous residue.

Step 2

$$O \longrightarrow OC_2H_5$$
 Br
 CF_3

9 g (39 mmol) of ethyl 2-methyl-4-trifluoromethyl-benzoate are dissolved in 200 ml of tetrachloromethane and admixed with 7 g (39 mmol) of *N*-bromo-succinimide and 0.1 g of dibenzoyl peroxide. The mixture is heated under reflux for 6 hours, and the precipitated succinimide is then filtered off and the filtrate is concentrated under water pump vacuum.

This gives 12 g of an amorphous residue which, in addition to ethyl 2-bromomethyl-4-trifluoromethyl-benzoate, contains 17% of ethyl 2,2-dibromomethyl-4-trifluoromethyl-benzoate and 12% of ethyl 2-methyl-4-trifluoromethyl-benzoate.

Step 3

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4 g of ethyl 2-bromomethyl-4-trifluoromethyl-benzoate (approximately 70% pure) and 2.28 g (12.8 mmol) of 5-bromo-4-methyl-2,4-dihydro-3H-1,2,4-triazol-3-one are dissolved in 150 ml of acetonitrile, admixed with 5.3 g (38.4 mmol) of potasssium carbonate and heated under reflux with vigoruous stirring for 2 hours. The reaction mixture is taken up in water and extracted repeatedly with methylene chloride. The combined methylene chloride phases are dried over sodium sulphate, concentrated under water pump vacuum and chromatographed.

This gives 2 g (38% of theory) of 5-bromo-4-methyl-2-(2-ethoxycarbonyl-5-trifluoromethyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one as an amorphous product. 1 H-NMR (CDCl₃, δ): 5.46 ppm.

Example (IV-2)

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6.7 g (40 mmol) of 4-methyl-5-trifluoromethyl-2,4-dihydro-3H-1,2,4-triazol-3-one are initially charged in 150 ml of acetonitrile and admixed with 11 g (80 mmol) of potassium carbonate. The mixture is heated to 50°C, and a solution of 13.1 g (44 mmol) of methyl 3-bromomethyl-2,4-dichloro-benzoate in 20 ml of acetonitrile is then added dropwise with stirring, and the reaction mixture is heated under reflux with stirring for another 15 hours. The mixture is subsequently concentrated under water pump vacuum, and the residue is taken up in methylene chloride, washed with 1N hydrochloric acid, dried with sodium sulphate and filtered. The filtrate is concentrated under reduced pressure, the residue is digested with petroleum ether and the crystalline product is isolated by filtration with suction.

This gives 14.9 g (97% of theory) of 4-methyl-5-trifluoromethyl-2-(2,6-dichloro-3-methoxycarbonyl-benzyl)-2,4-dihydro-3H-1,2,4-triazol-3-one of melting point 109°C.

By the methods of Examples (IV-1) and (IV-2), it is also possible to prepare. for example, the compounds of the general formula (IVa) listed in Table 3 below.

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$$X \xrightarrow{O} (R^4)_n (IVa)$$

$$X \xrightarrow{1} A Z$$

Table 3: Examples of compounds of the formula (IV)

Ex.	(position-)	(position-)	(position-)		physical data
No.	R^3	$(R^4)_n$	-A-Z	X	
IV-3	(2-) Cl	(4-) Cl	01	OCH_3	m.p.: 229°C
			N—CH ₃		$logP = 2.27^{a}$
			(3-) SCH ₃		
IV-4	(2-) Cl	(4-) Cl	O	OCH ₃	m.p.: 120°C
			N—CH ₃		$logP = 2.38^{a}$
			OC ₂ H ₅		
IV-5	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 127°C
					$logP = 2.55^{a}$
IV-6	(2-) Cl	(4-) Cl	O	OCH,	m.p.: 121°C
			$\begin{array}{c c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$		$logP = 2.04^{at}$

Ex.	(position-)	(position-)	(position-)	X 7	physical data
No.	R ³	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	X	
IV-7	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 68°C
			O		$\log P = 2.73^{\text{ a)}}$
			N/CH3		
			N CH ₃		
			OC ₃ H ₇ -i		
IV-8	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 129°C
			O.		$logP = 2.72^{a}$
			N CH ₃		
			N N N N N N N N N N N N N N N N N N N		
			OCH ₂ CF ₃		
			00112013		
IV-9	(2-) Cl	(4-) Cl	O	OCH ₃	m.p.: 164°C
			N CH ₃		$logP = 2.18^{a}$
			Br		·
			(3-)		
IV-10	(2-) Cl	(4-) Cl		OCH ₃	m.p.: 158°C
			N CH ₃		$logP = 1.55^{a}$
			N=		
			(3-) H		
			(3-)	OCH	10000
IV-11	(2-) Cl	(4-) Cl		OCH ₃	m.p.: 106°C
			N CH ₃		$\log P = 2.16^{a}$
			N=(
			(3-)	1	
137.10	(2) (1)	(4) (1	(3-)	OCH ₃	m.p.: 126°C
IV-12	(2-) Cl	(4-) Cl	(3-)	00113	
					$logP = 2.11^{a}$
			N CH ₃		
			N=(
			N(CH ₃) ₂		

Ex.	(position-)	(position-)	(position-)	v	physical data
No.	R ³	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	X	146°C
IV-13	(2-) Cl	(4-) Cl		OCH_3	m.p.: 146°C
			N CH ₃		$logP = 1.65^{a}$
			N 3		
			N		
			(3-) CH ₃		
				OCH	170°C
IV-14	(2-) Cl	(4-) Cl		OCH ₃	m.p.: 178°C
					$logP = 1.86^{a}$
			N		
			(3-)		
IV-15	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 97°C
14-13	(2-) (1	(,) (,			$\log P = 2.36^{a}$
					$\log 1 - 2.50$
			N N		
			\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
			N—		
			,OCH ³		
IV-16	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 99°C
10-10	(2-) C1	(4-) 61			$\log P = 2.73^{a}$
					$\log P = 2.73$
			\\\\\\\\\\		
			N		
1			OC ₂ H ₅		
TT 1.17	(2.) (1	(4-) Cl	(3-)	OCH ₃	m.p.: 56°C
IV-17	(2-) Cl	(4-) C1	(3-)	,	
					$logP = 3.08^{a}$
			N N		
	10	İ	N=		
			OC ₃ H ₇ -i		
		(1) (2)	(2)	OCH	m.p.: 102°C
IV-18	3 (2-) Cl	(4-) Cl	(3-)		
			O .		$\log P = 3.05^{a}$
1			N N		
			N=(
			OCH ₂ CF ₃		

Ex.	(position-)	(position-)	(position-)		physical data
No.	R^3	$(\mathbf{R}^4)_{\rm n}$	-A-Z	X	
IV-19	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 131°C
			N N		$logP = 2.70^{a}$
IV-20	(2-) Cl	(4-) Cl	SCH ₃	OCH ₃	m.p.: 135°C logP = 1.97 a)
			CH ₃		
IV-21	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 143°C
			N N N		$logP = 2.42^{a}$
ļ t			N(CH ₃) ₂		
IV-22	(2-) Cl	(4-) Cl	(3-)	OCH ₃	m.p.: 85°C
11-22	(2-) CI	(4-) CI	N OC ₂ H ₅		$\log P = 2.58^{a}$
			C_2H_5		
IV-23	(2-) Cl	(4-) Cl	N CH _s	OCH ₃	$logP = 1.98^{a}$
			(3-)		
IV-24	(2-) Cl	(4-) Cl	(3-)	OCH ₃	$logP = 2.07^{a}$
			N CH ₃		

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	
IV-25	(2-) Cl	(4-) Cl	0,	OCH ₃	m.p.: 157°C
					$logP = 2.94^{a}$
			SO ₂		
			(3-)		
IV-26	(4-) CF ₃	Н	(2-)	OC_2H_5	¹H NMR
			Q		(CDCl ₃ , δ):
			N CH ₃		5.53 ppm.
			SO ₂ CH ₃		
			2 3	OCH	TAL NIN (I)
IV-27	(4-) NO ₂	Н		OC ₂ H ₅	'H NMR
			N CH3		(CDCl ₃ , δ):
			N=<		5.48 ppm.
			(3-) CF ₃		
IV-28	(4-) NO ₂	Н	(3-)	OC_2H_5	¹H NMR
1 4 - 2 6	(4-)1102			0 02-3	(CDCl ₃ , δ):
			Ĭ		
			N N		5.30 ppm.
			N=<		
ļ					
IV-29	(4-)	Н	0	OC ₂ H ₅	'H NMR
	SO ₂ CH ₃		N CH ₃		(CDCl ₃ , δ):
			N=(5.61 ppm.
	1	!	CF		* *
			(3-)		
IV-30	(4-) Cl	Н	O	OC ₂ H ₅	
			N CH ₃		$(CDCl_3, \delta)$:
			N=		5.08 ppm.
			(3-) CH ₃		

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	X	IVI NO
IV-31	(4-) Cl	Н	O	OC_2H_5	¹H NMR
			N CH ₃		(CDCl ₃ , δ):
			N=(5.17 ppm.
			CF ₃		
			(3-)	00.11	ITTNINAD
IV-32	(4-) Cl	Н	(3-)	OC ₂ H ₅	¹H NMR
			O A		$(CDCl_3, \delta)$:
			\sim N \sim N \sim		5.00 ppm
			"		
137.22	(1)	Н	0	OC ₂ H ₅	$logP = 1.53^{a}$
IV-33	(4-)	11			
	SO ₂ CH ₃		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
			(2-)		
IV-34	(4-) Br	H	(2-)	OC ₂ H ₅	$\log P = 3.24^{\text{ a}}$
			Q		
			a OC H		
			NOC ₂ H ₅		
			$N = C_2H_5$		
			2.15		1 D 2 40 a)
IV-35	(4-) Br	Н		OC ₂ H	$\log P = 3.40^{a}$
			N N CH	3	
			N=(
			`CF.		
			(2-)	OCI	$1 \log P = 2.41^{a}$
IV-30	6 (4-) F	Н			$\log P = 2.41^{a}$
			N N CH	3	
) N=		
			(3-) Br		

Ex.	(position-)	(position-)	(position-)		physical data
No.	R^3	$(R^4)_n$	-A-Z	X	
IV-37	(4-) F	Н	N CH ₃	OC ₂ H ₅	$logP = 2.45^{a}$
IV-38	(4-) Br	H	0	OC ₂ H ₅	$logP = 2.06^{a}$
			$(3-)$ $ \begin{array}{c} N \longrightarrow CH_3 \\ CH_3 \end{array} $		
IV-39	(4-) Br	Н	0 N CH ₃ Br		$logP = 2.64^{a}$
IV-40	(4-) Br	Н	O N—CH ₃ CF ₃	OC ₂ H ₅	$logP = 3.23^{a}$
IV-41	(4-) Br	H	(3-) N	OC ₂ H ₅	$log P = 3.02^{a}$
IV-42	(4-) Cl	Н	O N N O	OC ₂ H ₂	$\log P = 3.23^{(a)}$

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	2 21 8
IV-43	(4-) Cl	Н	N=CH ₃	OC ₂ H ₅	$logP = 3.31^{a}$
			(2-)		
IV-44	(4-) Cl	Н	(2-)	OC ₂ H ₅	$logP = 3.14^{-a}$
			$ \begin{array}{c c} & \\ & \\ & \\ & \\ & \\ & \\ & \\ & $		
IV-45	(4-) NO ₂	Н	(2-)	OC ₂ H ₅	$logP = 2.42^{a}$
			N N N N N N N N N N		
IV-46	(4-) NO ₂	Н	N CH ₃ SCH ₃	4	$\log P = 2.82^{a}$
IV-47	7 (4-) CF ₃	H	(2-)	OC_2H	$\log P = 3.48^{a}$
			O N O		
IV-4	8 (4-) CF ₃	Н	(2-) N OC ₂ H ₅	OC ₂ F	$H_5 \log P = 3.38^{a}$
			$ \begin{array}{c} $		

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	
IV-49	(4-) CF ₃	H	N CH ₃	OC ₂ H ₅	$logP = 3.02^{a}$
			(2-) SCH ₃	OC ₃ H ₇	$logP = 3.91^{a}$
IV-50	(4-) CF ₃	Н	$\begin{array}{c c} & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ &$		logr = 3.91
IV-51	(4-) OCH ₃	Н	N CH ₃ N=(2-)	OC ₂ H ₅	
IV-52	(4-) OCH ₃	Н	$\begin{array}{c} O \\ N \\ N \end{array} \begin{array}{c} O \\ O \\ C_2 H_5 \end{array}$	OC ₂ H ₅	
IV-53	(4-) CF ₃	Н	N N N CH_3 OC_2H_5	OC ₂ H ₅	(CDCl ₃ , δ): 5.37 ppm.
IV-54	(4-) CF ₃	Н	$ \begin{array}{c} $	OC ₂ H ₅	'H NMR (CDCl ₃ , δ): 5.37 ppm.

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbf{R}^3	$(R^4)_n$	-A-Z	X	
IV-55	Н	Н	N CH ₃	OC ₂ H ₅	
IV-56	Н	Н	(2-) OC ₂ H ₅	OC ₂ H ₅	¹H NMR (CDCl ₃ , δ):
			(2-) OCH ₃	OC II	5.37 ppm.
IV-57	Н	Н	$\begin{array}{c c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$	OC ₂ H ₅	(CDCl ₃ , δ): 5.40 ppm.
IV-58	(4-) Br	Н	$N = \begin{pmatrix} O \\ N - CH_3 \end{pmatrix}$ OC_2H_5	OC ₂ H ₅	$logP = 2.95^{a}$
IV-59	(4-) Br	Н	0 N CH ₃ OCH ₃	OC ₂ H ₅	(CDCl ₃ , δ): 5.31 ppm.
IV-60	(4-) Br	Н	(2-)	OC ₂ H ₅	$logP = 2.44^{a}$

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{\mathbf{n}}$	-A-Z	X	LILNIMD
IV-61	(4-) F	Н		OC ₂ H ₅	¹H NMR
			N N CH ₃		(CDCl ₃ , δ):
			N=(5.35 ppm.
			OC ₂ H ₅		
*** 60	-4 · E	TY	0	OC_2H_5	¹H NMR
IV-62	(4-) F	Н	Ĭ	002115	i
			N CH ₃		(CDCl ₃ , δ):
			N=(5.53 ppm.
			(2-) CF ₃		
IV-63	(4-) F	Н	(2-)	OC_2H_5	¹H NMR
i			Q		(CDCl ₃ , δ):
			$N \longrightarrow N \longrightarrow OC_2H_5$		5.40 ppm.
			C ₂ H ₅		
IV-64	(4-) F	Н	0	OC_2H_5	¹H NMR
1 V - 04	(4-) 1		Ĭ cu	0 0 2 2 2 3	$(CDCl_3, \delta)$:
			N CH ₃		
			N=OCH ₃		5.36 ppm.
			(2-)		
IV-65	(4-) Br	Н	(2-)	OC ₂ H ₅	$\log P = 3.34^{a}$
			O		
			N CH ₃		
			N N O 1 13		
			OC ₃ H ₇ -i		
IV-66	(4-) Br	Н	(2-)	OC ₂ H ₅	$\log P = 3.38^{a}$
			O		
			N CH ₃		
			N=(
			OC ₃ H ₇ -n		

Ex.	(position-)	(position-)	(position~)		physical data
No.	\mathbb{R}^3	$(\mathbf{R}^4)_{n}$	-A-Z	X	
IV-67	(4-) Br	Н	(2-) N CH ₃	OC ₂ H ₅	$logP = 3.31^{a}$
IV-68	(4-) Br	Н	OCH ₂ CF ₃	OC ₂ H ₅	$logP = 2.16^{a}$
IV-69	(4-) Br	H	(2-) N=	OC ₂ H ₅	$logP = 2.41^{a}$
107	(1)		N N N N N N N N N N N N N N N N N N N		
IV-70	(4-) CF ₃	Н	N N N N N N N N N N	OC ₂ H ₅	$logP = 3.51^{a}$
IV-71	(4-) CF ₃	Н	(2-) N CH ₃ N OC ₃ H ₇ -n	OC ₂ H ₅	$logP = 3.54^{a}$
IV-72	(4-) Br	Н	N 0	OC ₂ H ₅	$\log P = 2.36^{a}$

Ex.	(position-)	(position-)	(position-)		physical data
No.	R^3	$(R^4)_n$	-A-Z	X	
IV-73	(4-) Br	Н	N O CH ₃	OC ₂ H ₅	$logP = 2.88^{a}$
IV-74	(4-) CF ₃	Н	(2-) N CH ₃		$logP = 2.68^{a}$
IV-75	(4-) Br	Н	N—CH ₃ N=(2-)		$\log P = 2.80^{a}$
IV-76	(4-) CF ₃	Н	(3-) N N OCH ₃		$logP = 3.87^{a}$
IV-77	(4-) CF ₃	Н	0 N N CH ₂ OCH ₃		$logP = 2.88^{a}$
IV-78	(4-) CF ₃	Н	(2-) N— CH ₂ OCH ₃	OC ₂ H ₅	$logP = 2.60^{a}$

Ex.	(position-)	(position-)	(position-)	W	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	$logP = 3.35^{a}$
IV-79	(4-) CF ₃	Н	(2-) N	OC ₂ H ₅	logr = 3.33
			Br		
IV-80	(4-) Br	Н	N CH ₃ SCH ₃	OC ₂ H ₅	$logP = 2.86^{a}$
IV-81	(4-) Cl	Н	0	OC ₂ H ₅	$logP = 2.83^{a}$
1 V - 81	(4-) CI	11	N—CH ₃		
			(2-)		
IV-82	(4-) Br	Н	(2-) N CH ₃ N(CH ₃) ₂	OC ₂ H ₅	$logP = 2.60^{a}$
111.02	14 × CE	7.7		OC ₂ H ₅	¹H NMR
IV-83	(4-) CF ₃	Н	$ \begin{array}{c c} & O \\ & N \\$		(CDCl ₃ , δ): 5.36 ppm.
			OC ₂ H ₅		
IV-84	(4-) CF ₃	Н	(2-)	OC ₂ H ₅	•
			$ \begin{array}{c} O \\ N \\ \longrightarrow \\ OCH_3 \end{array} $		(CDCl ₃ , δ): 5.37 ppm.

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	\mathbf{X}_{-}	
IV-85	(4-) CF ₃	Н	(2-)	OC_2H_5	$logP = 2.79^{a}$
			0		
			Ĭ		
!			N CH ₃		
) N=		
			N(CH ₃) ₂		
			. 0.2		0 (5 3)
IV-86	(4-) CF ₃	Н	Q'	OC ₂ H ₅	$logP = 3.67^{a}$
			/-N		
			$(2-)$ so_2		
				OC H	$logP = 3.80^{a}$
IV-87	(4-) CF ₃	Н		$OC_2\Pi_5$	log1 = 3.80
			(2-)		
137.00	(3-) CH ₃	Н	0	OC ₂ H ₅	$\log P = 2.54^{\text{ a}}$
IV-88	(3-) Cn ₃	11		2	
			N CH ₃		
			$N=\langle$		
			OC ₂ H ₅		
			(2-)		
IV-89	(4-)	Н	O	OC ₂ H ₅	$logP = 1.82^{a}$
			N CH ₃		
	SO ₂ CH ₃	Ì	N N 3		
			N=		
			(2-) SCH ₃		
			\\\-\\\-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	OC II	$\log P = 2.93^{\text{a}}$
IV-90	$(4-) CF_3$	Н		UC ₂ H ₂	logr - 2.93
			N		
1					
			CF ₃		
			(2-)		

Ex.	(position-)	(position-)	(position-)		physical data
No.	\mathbf{R}^3	$(R^4)_n$	-A-Z	X	
IV-91	(4-) CF ₃	Н	(2-)	OC ₂ H ₅	$logP = 3.08^{a}$
			N N		
			осн ₃		
IV-92	(4-) CF ₃	Н		OC ₂ H ₅	$logP = 3.04^{a}$
			N=C	i	
***	(1) CF	11	(2-) CH ₃	OC ₂ H ₅	$\log P = 3.45^{a}$
IV-93	(4-) CF ₃	Н	(2-)	002115	log1 3.43
			N CH ₃		
			OCH ₂ CF ₃		
IV-94	(4-) F	Н	(2-) O	OC ₂ H ₅	$\log P = 2.21^{a}$
			N CH ₃		
			N— N(CH ₃) ₂		
IV-95	(4-) F	Н	(2-)	OC ₂ H ₅	$\log P = 2.96^{a}$
			N CH ₃		
			OC ₃ H ₇ -n		
IV-96	(4-) F	Н	(2-) Q	OC ₂ H ₅	$logP = 2.05^{a}$
			N CH ₃		
			CH ₂ OCH ₃		

Ex.	(position-)	(position-)	(position-)	**	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	1 D 2 50 a)
IV-97	(4-) F	Н	(2-) O N	OC_2H_5	$logP = 2.50^{a}$
			OCH ₃		2 00 3
IV-98	(4-) F	Н	(2-)	OC_2H_5	$logP = 2.89^{a}$
			$ \begin{array}{c} O \\ N \\ O \\ O \\ O \\ O \\ O \\ O \\ O \\ O \\ O \\ O$		
IV-99	(4-) CF ₃	Н	O CH ₃ CH ₃ CH ₃	OC ₂ H ₅	$logP = 2.91^{-a}$
			(2-)		
IV-	(4-) Cl	Н	O _{II}	OC_2H_5	¹H NMR
100			N CH ₃		(CDCl ₃ , δ):
			$N = \langle$		5.39 ppm.
			(2-) CH ₃		
IV-	(4-) Cl	Н	O _{II}	OC ₂ H ₅	'H NMR
101			N CH ₃		$(CDCl_3, \delta)$:
			N=		5.50 ppm.
			(2-) CF ₃		
IV-	(4-) Cl	Н	(2-)	OC ₂ H ₅	¹H NMR
102			Ö		(CDCl ₃ , δ):
			N CH ₃		5.49 ppm.
			SO ₂ CH ₃		

Ex.	(position-) R ³	(position-)	(position-) -A-Z	X	physical data
No. IV-	(4-) CF ₃	(R ⁴) _n	-A-L	OC_2H_5	¹H NMR
	(4-) C13			2 3	$(CDCl_3, \delta)$:
103			N N CH ₃		_
			N=(CI		5.29 ppm.
			(2-) CH ₃		
IV-	(4-) CF ₃	Н	Q	OC ₂ H ₅	¹H NMR
104			N CH ₃		(CDCl ₃ , δ):
			N=(5.53 ppm.
			(2-) CF ₃		
77.7	(4.) CE	Н	(2-)	OC_2H_5	¹H NMR
IV-	(4-) CF ₃	H	(2-)	002115	
105					(CDCl ₃ , δ):
			\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		5.34 ppm.
			N=(
***		TT	(2)	OC ₂ H ₅	¹H NMR
IV-	(4-)	Н	(2-)	002115	
106	SO ₂ CH ₃				(CDCl ₃ , δ):
			N N		5.39 ppm.
			N=(
				OCII	LINIMD
IV-	(4-)	Н		OC ₂ H ₅	'H NMR
107	SO ₂ CH ₃		N N CH ₃		(CDCl ₃ , δ):
			N=		5.43 ppm.
		!	(2-) CH ₃		
IV-	(4-)	Н	(2-)	OC ₂ H ₅	¹H NMR
108	SO ₂ CH ₃		Q		$(CDCl_3, \delta)$:
			N CH ₃		5.40 ppm.
			N=\N>0.13		
			N(CH ₃) ₂		
			, 5, 2		

Ex.	(position-)	(position-)	(position-)	v	physical data
No.	R ³	$(R^4)_n$	-A-Z	X	III NIMD
IV-	(4-)	Н		OC_2H_5	¹H NMR
109	SO ₂ CH ₃		N CH ₃		(CDCl ₃ , δ):
			N=(5.38 ppm.
			(2-) OC ₂ H ₅		
IV-	(4-) Br	Н	Q	OC ₂ H ₅	¹H NMR
110			N CH ₃		(CDCl ₃ , δ):
					5.49 ppm.
			CF ₃		Provide the second
			(2-)		
IV-	Н	Н	(2-)	OC_2H_5	¹H NMR
111					(CDCl ₃ , δ):
			N		5.3 ppm.
			N=(
					ļ
IV-	Н	Н	O	OC ₂ H ₅	¹H NMR
112			N CH ₃		$(CDCl_3, \delta)$:
			N=(5.44 ppm.
			SCH ₃		
			(2-)		1 5 0 50 3)
IV-	(4-) CF ₃	Н	0	OC ₂ H ₅	$\log P = 2.58^{a}$
113			N CH ₃		
			N-		
			H_3C	1	
IV-	(4-)	Н	0	OCH,	$\log P = 1.53^{a}$
	Ļ		N CH ₃		
114	SO ₂ CH ₃		N N OII3		1
			N SCH ₃		
			(2-)		

Ex.	(position-)	(position-)	(position-)	-	physical data
No.	\mathbb{R}^3	$(R^4)_n$	-A-Z	X	
IV-	(4-)	Н	O II	OCH_3	$logP = 1.59^{a}$
115	SO ₂ CH ₃		N CH ₃		
	, 2-2-3				
			OC-H-		
			(2-)		
IV-	(4-) I	Н	O	OCH ₃	$logP = 2.68^{a}$
116			N CH ₃		
			OC_2H_5		
			(2-)		
IV-	(4-) CF ₃	Н	O _{II}	OCH ₃	$\log P = 2.74^{\text{ a}}$
117			N CH ₃		
			$N = \begin{pmatrix} 1 & 1 & 1 \\ 1 & 1 & 1 \end{pmatrix}$		
			OC ₂ H ₅		
			(2-)		
IV-	(4-) CF ₃	Н	0	OCH ₃	$logP = 2.65^{a}$
118			N CH ₃		
			$N = \langle 1 \rangle$		
			SCH ₃		
			(2-)		
IV-	(4-) CF ₃	H	O	OC_2H_5	$\log P = 2.96^{a}$
119			N CH ₃		
			N=(
			(2-) Br		
			(2-)	OCII	106°C
IV-	Н	Н		OCH ₃	m.p.: 106°C
120			N N CH ₃		
			N-		
			(2-) H ₃ C O		
				OCH	$\log P = 2.27^{a}$
IV-	(2-) NO ₂	(3-) OCH ₃		OCH ₃	$\frac{10gr - 2.27}{1}$
121			N N CH ₃		
			N=		
			OC ₂ H ₅		
			()		

5

10

Ex.	(position-) R ³	(position-)	(position-)	v	physical data
No.	R	$(R^4)_n$	-A-Z	_X	
IV-	$(2-) NO_2$	(3-) OCH ₃	O I	OCH_3	$logP = 2.19^{a}$
122			N N N N N N N N N N		

The logP values given in the Tables were determined in accordance with EEC Directive 79/831 Annex V.A8 by HPLC (High Performance Liquid Chromatography) using a reversed-phase column (C 18). Temperature: 43°C.

- (a) Mobile phases for the determination in the acidic range: 0.1% aqueous phosphoric acid, acetonitrile; linear gradient from 10% acetonitrile to 90% acetonitrile the corresponding data in the Tables are labelled a).
- (b) Mobile phases for the determination in the neutral range: 0.01-molar aqueous phosphate buffer solution, acetonitrile; linear gradient from 10% acetonitrile to 90% acetonitrile the corresponding data in the Tables are labelled b).
- 15 Calibration was carried out using unbranched alkan-2-ones (having 3 to 16 carbon atoms) whose logP values are known (determination of the logP values using the retention times by linear interpolation between two successive alkanones).

The lambda-max values were determined using the UV spectra from 200 nm to 400 nm in the maxima of the chromatographic signals.

Use examples:

Example A

5 Pre-emergence test

Solvent:

5 parts by weight of acetone

Emulsifier:

1 part by weight of alkylaryl polyglycol ether

To produce a suitable preparation of active compound, I part by weight of active compound is mixed with the stated amount of solvent, the stated amount of emulsifier is added and the concentrate is diluted with water to the desired concentration.

Seeds of the test plants are sown in normal soil. After approximately 24 hours, the soil is sprayed with the preparation of active compound such that the particular amount of active compound desired is applied per unit area. The concentration of the spray liquor is chosen so that the particular amount of active compound desired is applied in 1000 litres of water per hectare.

20

After three weeks, the degree of damage to the plants is assesssed in % damage in comparison to the development of the untreated control.

The figures denote:

25

30

0% = no effect (like untreated control)

100% =

total destruction

In this test, for example the compounds of Preparation Example 1 and 10 exhibit strong activity against weeds, and some of them are tolerated well by crop plants, such as, for example, maize.

Table A: Pre emergence test/greenhouse

Abutilon	100	06
Maize Cyperus Abutilon	001	100
Maize	1	0
Amount used (g ai./ha)	1000	200
Active compound of Preparation Example No.	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	(10)

Example B

Post-emergence test

5 Solvent:

10

15

20

5 parts by weight of acetone

Emulsifier:

1 part by weight of alkylaryl polyglycol ether

To produce a suitable preparation of active compound, 1 part by weight of active compound is mixed with the stated amount of solvent, the stated amount of emulsifier is added and the concentrate is diluted with water to the desired concentration.

Test plants which have a height of 5 - 15 cm are sprayed with the preparation of active compound in such a way that the particular amounts of active compound desired are applied per unit area. The concentration of the spray liquor is chosen so that the particular amounts of active compound desired are applied in 1000 l of water/ha.

After three weeks, the degree of damage to the plants is assesssed in % damage in comparison to the development of the untreated control.

The figures denote:

0% = no effect (like untreated control)

100% = total destruction

25

In this test, for example the compounds of Preparation Example 10 and 15 exhibit strong activity against weeds, and some of them are tolerated well by crop plants. such as, for example, maize.

Post emergence test/greenhous
Table B:

Sinapis	08	80
Maize Amaranthus	95	06
Maize	20	0
Amount used (g ai./ha)	200	1000
Active compound of Preparation Example No.	$\begin{array}{c c} & & & & & & & & & & & & & & & & & & & $	(15)